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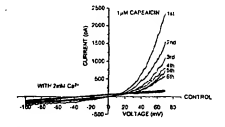
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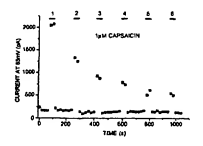
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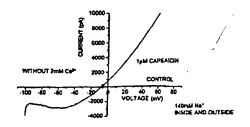
(54) Title: HUMAN VANILLOID RECEPTORS AND THEIR USES

(57) Abstract

The invention provides novel human vanilloid receptor (hVR) proteins, in particular hVR1 and hVR3, nucleotide sequences encoding for the novel hVR proteins, and hVR proteins for use in a method for screening for agents useful in the treatment or prophylaxis of disorders which are responsive to modulation of hVr activity in a human patient. The invention also provides expression vectors comprising said nucleotide sequences, stable cell lines comprising said expression vectors, antibodies specific for the novel hVR proteins, methods for the identification of compounds which exhibit hVR modulating activity, compounds identifiable and identified by such methods, and methods of treatment or prophylaxis of disorders which are responsive to modulation of hVR activity in a human patient.







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HUMAN VANILLOID RECEPTORS AND THEIR USES

Field of the Invention

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The present invention relates to human vanilloid receptor (hVR) proteins and to related nucleotide sequences, expression vectors, cell lines, antibodies screening methods, compounds, methods of production and methods of treatment, as well as other related aspects.

Background of the Invention

10 Capsaicin, the irritant in hot peppers and a member of the vanilloid family activates a sub-group of sensory neurons: the nociceptors. These neurons transmit nociceptive and thermoceptive pain information back to pain-processing centres in the central nervous system such as the spinal cord and the brain. They are also sites for the release of pro-inflammatory mediators in the periphery (1). Nociceptors show heterogeneity in their sensitivity to capsaicin. 15 Excitation and prolonged exposure of these neurons to capsaicin is followed by a refractory state known as desensitisation (2) when they become insensitive to capsaicin and other noxious stimuli (3). The long-term response to insensitivity could be explained by death of the nociceptors or destruction of its peripheral terminals (4). Because of the desensitisation phenomenon, capsaicin has been 20 used therapeutically for decades as an analgesic agent for the treatment of pain in a range of disorders (5).

It has been speculated that the endogenous target for capsaicin plays an important function in the detection of painful stimuli. It has been shown by electrophysiological and biochemical studies that capsaicin induces a flux of cations in dorsal root ganglion (DRG) neurons (6,7). Because other vanilloid derivatives show responses in a dose dependent manner (8,9) a receptor is the most likely candidate to explain the mechanism. Therefore, based on indirect evidence it has been anticipated that these actions of capsaicin (excitation / desensitisation) are mediated by a specific membrane-bound receptor named vanilloid receptor (10).

Evidence for the existence of a vanilloid receptor came from binding experiments with resiniferatoxin (RTX), a capsaicin analog (11), and a competitive antagonist

of capsaicin, capsazepine (12). Vanilloid receptors have been visualised by using ([³H]-RTX) autoradiography in dorsal root ganglia (DRG) and spinal cord of different species including man (13,14).

Recently, a rat vanilloid receptor termed VR1 has been identified using an expression-cloning strategy to isolate the complementary DNA (cDNA) encoding the corresponding protein from a rat DRG cDNA library (15). The cDNA clone was completely sequenced. The rat VR1 cDNA has an open reading frame of 2,514 nucleotides and encodes for a protein of 838 amino acids with a predicted relative molecular mass of 95,000. Analysis of the amino acid sequence identified 6 potential transmembrane regions with a short hydrophobic stretch between the transmembrane regions 5 and 6. The N-terminus (amino terminal) contains three ankyrin repeat domains. No motifs have been identified at the C-terminus (carboxy terminal).

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It has been noted that rat VR1 transfected cells exhibit an increase in calcium levels after heat treatment and it has been suggested that *in vivo* VR1 and vanilloid receptors are involved in detection of noxious heat (but not innocuous heat). It has also been proposed that protons could act as modulators of the vanilloid receptors (16, 17, 18).

While it has been recognised that the rat capsaicin receptor, VR1, is a member of the family of non-selective ion channels that are gated by ligands and that it is involved in pain sensation, the natural ligand of VR1 remains unknown. It is therefore suggested that human vanilloid receptor sub-types may provide targets for the development of novel analgesic agents (agonists and antagonists) and agents which may interact with other disorders.

Accordingly, it is an object of the present invention to locate and characterise human vanilloid receptors. Other objects of the present invention will become apparent from the following detailed description thereof.

Summary of the Invention

According to one embodiment of the present invention there is provided an isolated human vanilloid receptor (hVR) protein or a variant thereof. Preferably

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the hVR protein is an hVR1 or hVR3 protein or a variant thereof. In a particularly preferred aspect of the invention the hVR protein has an amino acid sequence as shown in figure 3 or in figure 18.

According to another aspect of the invention, there is provided a human vanilliod receptor (hVR) protein or a variant thereof, preferably hVR1 or hVR3 or a variant thereof, for use in a method of screening for agents useful in the treatment or prophylaxis of a disorder which is responsive to the modulation of hVR activity, preferably hVR1 or hVR3 activity, in a human patient. Preferably the disorder is pain, neuropathic pain, inflammatory pain, chronic pain, post-operative pain, rheumatoid arthritic pain, neuropathies, neuralgia, algesia, neurodegeneration, nerve injury, stroke, ischaemia migraine, irritable bowl syndrome (IBS), a respiratory disorder such as asthma or chronic obstructive pulmonary disease (COPD), a urological disorder such as diabetic neuropathy, incontinence and interstitial cystitis, or an inflammatory disorder.

According to another aspect of the invention there is provided a nucleotide sequence encoding a human vanilloid receptor (hVR) protein or a variant thereof as hereinbefore described, or a nucleotide sequence that is complementary thereto. Preferably the nucleotide sequence encodes an hVR1, hVR3 protein or variant thereof or a nucleotide sequence which is complementary thereto. Particularly preferably the nucleotide sequence is as shown in figure 2 and figure 17.

According to another aspect of the invention there is provided an expression vector comprising a nucleic acid sequence as referred to above which is capable of expressing an hVR protein as hereinbefore described or a variant thereof, preferably hVR1 or hVR3 or a variant thereof. Preferably the expression vector is as displayed in figure 6 or figure 20.

According to another aspect of the invention there is provided a stable cell line comprising an expression vector as referred to above which is capable of expressing an hVR protein as hereinbefore described or a variant thereof, preferably hVR1 or hVR3 or a variant thereof. The stable cell line is preferably a

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modified mammalian cell line, preferably HEK293, CHO, COS, HeLa or BHK although transient expression may be preferred in *Xenopus* oocytes.

According to another aspect of the invention there is provided an antibody specific for an hVR protein as hereinbefore described or a variant thereof, preferably specific for hVR1 or hVR3 or a variant thereof.

According to another aspect of the invention there is provided a method for identification of a compound which exhibits hVR modulating activity, comprising contacting an hVR protein as hereinbefore described or a variant thereof, preferably hVR1 or hVR3 or a variant thereof, with a test compound and detecting modulating activity or inactivity.

According to another aspect of the invention there is provided a compound which modulates hVR activity, preferably that of hVR1 or hVR3, identifiable by the method referred to above.

According to another aspect of the invention there is provided a compound which modulates hVR activity, preferably that of hVR1 or hVR3, identifiable by the method referred to above, for use in therapy.

According to another aspect of the invention there is provided the use of a compound which modulates hVR activity, preferably that of hVR1 or hVR3, identifiable by the method referred to above, in the manufacture of a medicament for treatment or prophylaxis of a disorder which is responsive to the modulation of hVR activity, preferably hVR1 activity or hVR3 activity, in a human patient. Preferably the disorder is pain, neuropathic pain, inflammatory pain, chronic pain, post-operative pain, rheumatoid arthritic pain, neuropathies, neuralgia, algesia, neurodegeneration, nerve injury, stroke, ischaemia migraine, irritable bowl syndrome (IBS), a respiratory disorder such as asthma or chronic obstructive pulmonary disease (COPD), a urological disorder such as neuropathy, incontinence or interstitial cystitis, or an inflammatory disorder.

According to another aspect of the invention there is provided a method of treatment or prophylaxis of a disorder which is responsive to modulation of hVR,

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preferably hVR1 or hVR3, activity in a human patient which comprises administering to said patient an effective amount of a compound identifiable by the method referred to above. Preferably the disorder is pain, neuropathic pain, inflammatory pain, chronic pain, post-operative pain, rheumatoid arthritic pain, neuropathies, neuralgia, algesia, neurodegeneration, nerve injury, stroke, ischaemia migraine, irritable bowl syndrome (IBS), respiratory disorders such as asthma and chronic obstructive pulmonary disease (COPD) and urological disorders including diabetic neuropathy, incontinence and interstitial cystitis and inflammatory disorders.

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According to another aspect of the invention there is provided a compound which modulates hVR activity, preferably that of hVR1 or hVR3, identifiable by the method referred to above, excluding the compounds capsaicin, resiniferatoxin, piperine, zingerone, polydodial, warburganal, aframodial, cinnamodial, cinnamosmolide, cinnamolide, isovelleral, scalaradial, ancistrodial, β-acaridial, scutigeral, merulidial, anandamide and capsazepine.

According to another aspect of the invention there is provided a compound

which modulates hVR activity, preferably that of hVR1 or hVR3, identifiable by the method referred to above, excluding the compounds capsaicin, resiniferatoxin, piperine, zingerone, polydodial, warburganal, aframodial, cinnamodial, cinnamosmolide, cinnamolide, isovelleral, scalaradial, ancistrodial, β-acaridial, scutigeral, merulidial, anandamide and capsazepine, for use in

therapy.

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According to another aspect of the invention there is provided the use of a compound which modulates hVR activity, preferably that of hVR1 or hVR3, identifiable by the method referred to above, excluding the compounds resiniferatoxin, piperine, zingerone, polydodial, warburganal, aframodial, cinnamodial, cinnamosmolide, cinnamolide, isovelleral, scalaradial, ancistrodial, β-acaridial, scutigeral, merulidial, anandamide and capsazepine, in the manufacture of a medicament for treatment or prophylaxis of a disorder which is responsive to the modulation of hVR activity, preferably hVR1 activity or hVR3 activity, in a human patient. Preferably the disorder is pain, neuropathic pain, inflammatory pain, chronic pain, post-operative pain, rheumatoid arthritic

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pain, neuropathies, neuralgia, algesia, neurodegeneration, nerve injury, stroke, ischaemia migraine, irritable bowl syndrome (IBS), a respiratory disorder such as asthma or chronic obstructive pulmonary disease (COPD), a urological disorder such as neuropathy, incontinence or interstitial cystitis, or an inflammatory disorder.

According to another aspect of the invention there is provided a method of treatment or prophylaxis of a disorder which is responsive to modulation of hVR, preferably hVR1 or hVR3, activity in a human patient which comprises administering to said patient an effective amount of a compound identifiable by the method referred to above, excluding the compounds capsaicin, resiniferatoxin, piperine, zingerone, polydodial, warburganal, aframodial, cinnamodial, cinnamosmolide, cinnamolide, isovelleral, scalaradial, ancistrodial, β-acaridial, scutigeral, merulidial, anandamide and capsazepine. Preferably the disorder is pain, neuropathic pain, inflammatory pain, chronic pain, post-operative pain, rheumatoid arthritic pain, neuropathies, neuralgia, algesia, neurodegeneration, nerve injury, stroke, ischaemia migraine, irritable bowl syndrome (IBS), respiratory disorders such as asthma and chronic obstructive pulmonary disease (COPD) and urological disorders including diabetic neuropathy, incontinence and interstitial cystitis and inflammatory disorders.

According to another aspect of the invention there is provided a compound identified by the method referred to above.

According to another aspect of the invention there is provided a compound identified by the method referred to above, for use in therapy.

According to another aspect of the invention there is provided the use of a compound identified by the method referred to above in the manufacture of a medicament for treatment or prophylaxis of a disorder which is responsive to the modulation of hVR activity, preferably hVR1 activity or hVR3 activity, in a human patient. Preferably the disorder is pain, neuropathic pain, inflammatory pain, chronic pain, post-operative pain, rheumatoid arthritic pain, neuropathies, neuralgia, algesia, neurodegeneration, nerve injury, stroke, ischaemia migraine, irritable bowl syndrome (IBS), a respiratory disorder such as asthma or chronic

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obstructive pulmonary disease (COPD), a urological disorder such as neuropathy, incontinence or interstitial cystitis, or an inflammatory disorder.

According to another aspect of the invention there is provided a method of treatment or prophylaxis of a disorder which is responsive to modulation of hVR, preferably hVR1 or hVR3, activity in a human patient which comprises administering to said patient an effective amount of a compound identified by the method referred to above. Preferably the disorder is pain, neuropathic pain, inflammatory pain, chronic pain, post-operative pain, rheumatoid arthritic pain, neuropathies, neuralgia, algesia, neurodegeneration, nerve injury, stroke, ischaemia migraine, irritable bowl syndrome (IBS), respiratory disorders such as asthma and chronic obstructive pulmonary disease (COPD) and urological disorders including diabetic neuropathy, incontinence and interstitial cystitis and inflammatory disorders.

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According to another aspect of the invention there is provided a method of producing an hVR protein as hereinbefore described or a variant thereof, preferably hVR1 or hVR3 or a variant thereof, comprising introducing into an appropriate cell line a suitable vector comprising a nucleotide sequence encoding for an hVR protein or a variant thereof, preferably hVR1 or hVR3 or a variant thereof, under conditions suitable for obtaining expression of the hVR protein or a variant thereof, preferably hVR1 or hVR3 or a variant thereof.

Brief Description of the figures

25 Figure 1 is an alignment of hVR1 in silico derived clusters with rat VR1.

Figure 2 displays the human VR1 nucleotide sequence including the 5'UTR (nt – 773 to nt 0), coding region (nt 1 to 2517) and 3'UTR (nt 2518 to nt 3560).

Figure 3 illustrates the nucleotide and encoded amino acid sequence of the human VR1sequence.

Figure 4 depicts the amino acid sequence of the hVR1 gene, the shading denotes predicted trans-membrane regions (boxed) and the ankyrin binding domains (unboxed). The predicted phosphorylation sites are underlined.

Figure 5 is a comparison of the amino acid sequences of the rat (rVR1) and human (hVR1) vanilloid receptors.

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Figure 6 illustrates constructs pBluescriptSK(+) (A) and pCIN5-new (B) with the full length hVR1 gene cloned via Notl and EcoRI restriction sites.

Figure 7 shows a Slot Blot hybridisation with hVR1 probe with positive labelling of both rat and human DRG mRNA.

Figure 8 displays a Western blot probed with anti-VR1 antibodies with the arrow indicating the VR1 specific protein.

Figure 9 shows localisation of VR1 in rat DRG tissue sections, the arrow points to VR1 expressing small diameter ($<25\mu$ n) neurone cell bodies.

Figure 10 depicts the *in situ* localisation of VR1 in human DRG sections (A) and human skin (B).

Figure 11 illustrates the functional response to capsaicin and blockade by capsazepine (CPZ) (A) with the current voltage relationship plotted in (B) on human VR-1 channels, transiently expressed in HEK293T cells.

Figure 12 shows capsaicin-induced desensitisation of human VR-1 channels in the presence of 2mM external calcium (A), maximum current (65mV) against time (B) and current voltage relationship in the absense of Ca²⁺ (C).

Figure 13 shows the influx of calcium into transiently transfected HEK293T cells over a time course in the presence of agonist capsaicin, anandamide and resiniferatoxin in the absence (A, B, D and F) or presence (C, E, G) of the antagonist, capsezipine.

Figure 14 illustrates a graphical presentation the results shown in figure 13 examining the response of hVR1 transfected HEK293T cells over time before and after exposure to agonists: capsaicin, anandamide and resiniferatoxin in the absence (A, B, D and F) or presence (C, E, G) of the antagonist, capsezipine.

Figure 15 displays the proposed assay strategy to carry out drug screening. Figure 16 displays an alignment of *in silico* derived hVR3 specific clusters with rat VR1.

Figure 17 depicts the hVR3 nucleotide sequence including the 5' UTR (nt -686 to nt 0) Coding region (nt1 to nt 2889), 3'UTR (nt 2890 to nt 3418).

Figure 18 shows the nucleotide and amino acid sequence of hVR3.

Figure 19 is of the amino acid sequence of hVR3, the shading denotes predicted trans-membrane regions (boxed) and the ankyrin binding domains (unboxed).

Figure 20 displays constructs pBluescriptSK(+) (A) and pCDNA3.1 (+) (B) with the full length hVR3 gene cloned via NotI and XhoI restriction sites.

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Figure 21 illustrates a multiple comparison of the amino acid sequences of the rat VR1 and the human vanilloid receptors: hVR1, hVRL-1 and hVR3.

Figure 22 Northern Blot hybridisation with hVR3 probe with strong signals detected in trachea (A), prostate (B), placenta, kidney and pancreas (C).

Detailed Description of the Invention

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Throughout the present specification and the accompanying claims the words "comprise" and "include" and variations such as "comprises", "comprising", "includes" and "including" are to be interpreted inclusively. That is, these words are intended to convey the possible inclusion of other elements or integers not specifically recited, where the context allows.

As referred to above, the present invention relates to isolated human vanilloid receptor (hVR) proteins, and in particular to the human vanilloid receptors which will be termed respectively human vanilloid receptors 1 and 3 (hVR1, and hVR3), sequence information for which is provided in figures 2 (hVR1) and 17 (hVR3). In the context of this invention the term "isolated" is intended to convey that the receptor protein is not in its native state, insofar as it has been purified at least to some extent or has been synthetically produced, for example by recombinant methods. The term "isolated" therefore includes the possibility of the receptor protein being in combination with other biological or non-biological material, such as cells, suspensions of cells or cell fragments, proteins, peptides, organic or inorganic solvents, or other materials where appropriate, but excludes the situation where the receptor protein is in a state as found in nature.

Routine methods, as further explained in the subsequent experimental section, can be employed to purify and/or synthesise the receptor proteins according to the invention. Such methods are well understood by persons skilled in the art, and include techniques such as those disclosed in Sambrook, J. et al. (28), the disclosure of which is included herein in its entirety by way of reference.

By the term "variant" what is meant throughout the specification and claims is that other peptides or proteins which retain the same essential character of the human vanilloid receptor proteins for which sequence information is provided, are also intended to be included within the scope of the invention. For example,

other peptides or proteins with greater than about 80%, preferably at least 90% and particularly preferably at least 95% homology with the sequences provided are considered as variants of the receptor proteins. Such variants may include the deletion, modification or addition of single amino acids or groups of amino acids within the protein sequence, as long as the peptide maintains the biological functionality of a human vanilloid receptor. This biological functionality can of course be assessed by conducting binding studies with known vanilloid modulators such as capsaicin, capsazepine (12) and resiniferatoxin (11).

Human VR1 is preferentially expressed in human dorsal root ganglia (DRG) and relative to hVR3 has the highest sequence homology with the rat VR1. Therefore, hVR1 is likely to be the human orthologue to rat VR1. hVR3 is less similar to rat VR1 and is expressed in a wider range of tissues. Nucleotide sequence analysis of hVR1 reveals a 2517bp open reading frame which encodes an 839 amino acid protein (see figures 2, 3 and 4). This deduced hVR1 protein sequence is 86 % identical to the rat VR1 (15) and shares many of its characteristics such as 6 transmembrane regions with an hydrophobic stretch between transmembrane 5 and 6 and an N-terminus which contains 3 ankyrin repeat domains. Similarly hVR3 has an open reading frame of 2889bp open reading frame which encodes a 963 amino acid protein (see figures 17, 18 and 19). The deduced hVR3 protein is 46 % identical to rat VR1 and 44 % identical to hVR1 sharing many of VR1's characteristics such as 6 transmembrane regions with an hydrophobic stretch between transmembrane 5 and 6 and an N-terminus which contains 3 ankyrin repeat domains.

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The invention also includes nucleotide sequences which encode for human vanilloid receptor proteins or variants thereof as well as nucleotide sequences which are complementary thereto. Preferably the nucleotide sequence is a DNA sequence and most preferably, a cDNA sequence. Preferably the proteins are hVR1, hVR3 or variants thereof. Such nucleotides can be isolated or synthesised according to methods well know in the art. See reference 28, the disclosure of which is included herein in its entirety by way of reference.

The present invention also includes expression vectors which comprise nucleotide sequences encoding for the hVR, preferably hVR1 or hVR3, receptor

proteins or variants thereof. A further aspect of the invention relates to an expression vector comprising nucleotide sequences encoding for hVR1 or hVR3 receptor proteins or variants thereof. Such expression vectors are routinely constructed in the art of molecular biology and may for example involve the use of plasmid DNA and appropriate initiators, promoters, enhancers and other elements, such as for example polyadenylation signals which may be necessary, and which are positioned in the correct orientation, in order to allow for protein expression. Suitable vectors for use in practicing the present invention include pBluescript (Stratagene), pCR-Script (Stratagene), pCR2.1-TOPO (Invitrogen), pCRII-TOPO (Invitrogen), pCR-Blunt (Invitrogen), with vectors such as pCIN (32) (available from Clontech as pIRES-neo), pCDNA 3.1 (Invitrogen) or pClneo (Promega) required for mammalian expression. Appropriate methods can be effected by following protocols described in many standard laboratory manuals (28, 29).

The invention also includes cell lines which have been modified to express the novel receptor. Such cell lines include transient, or preferably stable higher eukaryotic cell lines, such as mammalian cells or insect cells, lower eukaryotic cells, such as yeast or prokaryotic cells such as bacterial cells. Particular examples of cells which have been modified by insertion of vectors encoding for the receptor proteins according to the invention include HEK293T cells and Xenopus oocytes. Preferably the cell line selected will be one which is not only stable, but also allows for mature glycosylation and cell surface expression of the inventive receptor. Representive examples of appropriate hosts include animal cells such as HEK293, CHO, COS, HeLa and BHK.

It is also possible for the receptors of the invention to be transiently expressed in a cell line or on a membrane, such as for example in a baculovirus expression system. Such systems, which are adapted to express the receptors according to the invention, are also included within the scope of the present invention.

In particular, the functional hVR protein may include hVR receptor proteins selected from hVR1 and hVR3 and thereof or even other hVR protein subtypes or splice variants which have not yet been identified.

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According to another aspect, the present invention also relates to antibodies, preferably monoclonal antibodies, which have been raised by standard techniques and are specific for the receptor proteins or variants thereof according to the invention. Such antibodies could for example, be useful in purification, isolation or screening involving immuno precipitation techniques and may be used as tools to further ellucidate hVR, preferably hVR1 or hVR3, protein function, or indeed as therapeutic agents in their own right. Antibodies may also be raised against specific epitopes of the receptors according to the invention.

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An important aspect of the present invention is the use of receptor proteins according to the invention in screening methods designed to identify compounds which act as receptor ligands and which may be useful to modulate receptor activity. In general terms, such screening methods will involve contacting the receptor protein concerned, preferably hVR1 or hVR3, with a test compound and then detecting modulation in the receptor activity, or indeed detecting receptor inactivity, which results. For further details on the screening strategy refer to figure 15. The present invention also includes within its scope those compounds which are identified as possessing useful hVR, preferably hVR1 or hVR3. modulation activity, by the screening methods referred to above. The screening methods comprehended by the invention are generally well known to persons skilled in the art. High throughput screens may include fluorescence based assays using the Fluorometric Imaging Plate Reader (FLIPR) with calcium sensitive dyes, and reporter gene assays using calcium sensitive photoproteins that emit light on the influx of calcium and can be detected using an Imaging system. Secondary screens may involve electrophysiological assays utilising patch clamp technology to identify small molecules; antibodies, peptides, proteins or other types of compounds that interact with hVR, preferably hVR1 or hVR3, to modulate activity. Tertiary screens may involve the study of modulators in well characterised rat and mouse models of pain. These models of pain include, but are not restricted to, intraplantar injection of inflammatory agents such as carageenan, formalin and complete freunds adjuvant (CFA). Models of neuropathic pain such as loose ligature of the sciatic nerve are also included. Other screens may involve the study of modulators in human volunteers subject to topically applied capsaicin.

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Another aspect of the present invention is the use of compounds which have been identified by screening techniques referred to above in the treatment or prophylaxis of disorders which are responsive to modulation of hVR, preferably hVR1 or hVR3, receptor activity, in a human patient. By the term "modulation" what is meant is that there will be either agonism or antagonism at the receptor site which results from ligand binding of the compound at the receptor. By the term "modulation" what is meant is that there will be either agonism or antagonism at the receptor site which results from ligand binding of the compound at the receptor excluding the compounds capsaicin, resiniferatoxin, piperine, zingerone, polydodial, warburganal, aframodial, cinnamosmolide, cinnamolide, isovelleral, scalaradial, ancistrodial, β-acaridial, scutigeral, merulidial, anandamide and capsazepine. hVR, preferably hVR1 and hVR3, proteins have been implicated in disorders of the central nervous system (CNS), gastrointestinal (GI) tract, lungs and bladder and therefore modulation of hVR, preferably hVR1 or hVR3, receptor activity in these tissues will result in a positive therapeutic outcome in relation to such disorders. In particular, the compounds which will be identified using the screening techniques according to the invention will have utility for treatment and/or prophylaxis of disorders such as pain, neuropathic pain, inflammatory pain, chronic pain, post-operative pain. rheumatoid arthritic pain, neuropathies, neuralgia, algesia, neurodegeneration, nerve injury, stroke, ischaemia migraine, IBS, respiratory disorders such as asthma and COPD, urological disorders including diabetic neuropathy, incontinence and interstitial cystitis, and inflammatory disorders. understood however, that the mention of such disorders is by way of example only, and is not intended to be limiting on the scope of the invention.

The compounds which are identified according to the screening methods outlined above may be formulated with standard pharmaceutically acceptable carriers and/or excipients as is routine in the pharmaceutical art, and as fully described in Remmington's Pharmaceutical Sciences, Mack Publishing Company, Eastern Pennsylvania, 17th Ed, 1985, the disclosure of which is included herein in its entirety by way of reference.

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PCT/EP99/09284

The compounds may be administered via enteral or parenteral routes such as via oral, buccal, anal, pulmonary, intravenous, intraarterial, intramuscular, intraperitoneal, topical or other appropriate administration routes.

The present invention will be further explained, by way of examples, in the appended experimental section. Reference examples are provided.

Experimental details

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10 Reference Example A: Identification of related human ESTs (Expressed Sequence Tags) (19) to the rat VR1 sequence by *in silico* analysis

The full-length rat VR1 amino acid sequence (15) was used as a query sequence using the tBlastn (20) alignment program to identify related human genes in the dbEST (21) and Incyte (Incyte Pharmaceuticals, Inc., 3174 Porter Drive, Palo Alto, California 94304, USA) databases. Several human ESTs were identified and those with similarities greater than 50% selected for further analysis. One of these ESTs was T12251 previously shown to have 68% aminoacid identity and 84% similarity over a region of 70 amino acids (15). Full-length cloning and functional characterisation of the gene represented by this cluster has been completed (30). This gene was denoted hVRL-1 and encoded a protein of 764 amino acid protein that was 48 % identical to the rat VR1 protein. All human ESTs from both databases were clustered to identify overlapping identical ESTs belonging to the same transcript. The GCG package (Wisconsin Package Version 9.0, Genetics Computer Group (GCG), Madison, Wisconsin) and a program developed in house termed ESTBlast (22) were used to build up these clusters. In total, forty-three ESTs derived from different tissue sources and both EST databases were clustered into ten groups, one of these clusters represented hVRL-1. The remaining nine clusters have been named hVRa, hVRb, hVRc, hVRd, hVRe, hVRf, hVRg, hVRh and hVRi. For each EST the tissue source was assigned according to the annotations in the dbEST and Incyte databases. Since no obvious starting codon was present and the cluster sequences were shorter than the rat VR1 transcript none of these clusters were likely to represent a full-length vanilloid receptor transcript. Finally hVRg, hVRh and hVRi collapsed into a single contig. Sequence analysis has shown that

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these cDNAs are likely to be chimeric. The 5' end has weak similarities with the rat VR1 gene but the 3' end is identical to a DNA binding protein. No more work was pursued with that transcript.

5 Reference Example B: Isolation of the human orthologue to the rat VR1 gene (reference examples B1-B4):

Reference Example B1: In silico assembly of human VR1

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The consensus nucleotide sequences from the ten clusters were searched with the tBlastx program (20) against the rat VR1 sequences to identify the most likely open reading frames. Frame shifts were corrected when the sequence trace files were available. Each cluster was aligned against the rat VR1 amino-acid sequence according to the Blastx results. The Blastx alignment program (20) was used to compare the full-length rat VR1 protein with the amino-acid sequences of the ten clusters. The three clusters with the highest homology, displayed in figure 1, were aligned with the rat VR1 gene.

Cluster hVRa shared a high homology (70% identity and 75% similarity over a stretch of 107 amino acids) with the 5' of the rat VR1 sequence but did not seem to have a potential start codon. It contained two ESTs (EST1 and EST2) derived from the same tissue, bladder, and from the same patient. These two ESTs were selected for further investigation since this cluster was the most 5', had high homology with rat VR1 and the bladder tissue could be contaminated with sensory neurones. Both cDNA clones were ordered but only clone EST1 was received as EST2 failed the recovery procedure.

Cluster hVRb composed of two EST's (EST3 and EST4), with 89% identity and 92% similarity over 90 residues, showed the highest degree of homology to the rodent sequence. The overlap between both sequences was located towards the middle of the gene.

hVRc (EST5) also while having high homology (71% identity and 75% similarity over 65 residues) with rat VR1 was closely related to the C-terminus of the rat protein sequence.

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Reference Example B2: Sequencing of clones

All DNA sequences were determined by automated DNA sequencing based on the dideoxy chain-termination method using the ABI 373A / 377 sequencers (Applied Biosystems). Sequence-specific primers were used with the 'Big-Dye' Terminator Cycle Sequencing kit (Applied Biosystems). The nucleotide sequence was analysed using programs from the University of Wisconsin Genetics Computer Group package.

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More specifically when sequencing an EST clone, the following protocol was adopted. The EST1 clone was grown using standard procedures and DNA was isolated using Qiagen columns. SP6 (5' ATTTAGGTGACACTATAG) and T7 (5' TAATACGACTCACTATAGGG) primers flanking the cloning site were used to sequence both ends. Plasmid DNA (0.6 pmol) was used with 10.0 pmol of each primer for the dye terminator reaction. The SP6 end corresponded to the *in silico* derived EST sequence (identical to EST1). The T7 end did not have homologies with VR1 nor did it possess a long open reading frame or a polyadenylation motif. The size of the insert was determined by enzyme digestion of the DNA with the endonucleases Notl and EcoRI and calculated to be approximately 3kb.

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Plasmid DNA (50ng) was used to amplify the insert by Polymerase Chain Reaction (PCR) with T7 and SP6 as primers. The PCR conditions included an initial hot-start at 94°C for 2 minutes, followed by 35 cycles at 94°C for 45 seconds, 50°C for 45 seconds and 72°C for 1 minute and terminated by 5 minutes at 72°C. The resulting PCR amplicon was separated on a 1.2% agarose gel and shown to be of ~3kb in size.

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To fully sequence the PCR product the nuclease-Bal-31 technique was used where both strands of duplex DNA are degraded from both ends (23). After ethanol precipitation of the PCR product, the pellet was re-suspended in 30ml of 1X Bal-31 buffer (add buffer composition). A time-course digest with 2 units of Bal-31 enzyme (Roche Molecular Biochemicals) was carried out with 12 time points taken over 90 minutes (30 seconds, 1, 2, 3, 5, 7, 10, 15, 25, 45, 75 and 90 minutes). Three pools were made respectively from digests 1 to 4, 5 to 8 and 9

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to 12. Each pool was blunt-ended and sub-cloned into the pCR-Script SK (+) plasmid from Stratagene at the Srfl site. After transformation, 16 colonies from each pool were screened by PCR with the flanking Reverse (5' GGAAACAGCTATGACCATG) and M13-20 (5' GTAAAACGACGGCCAGT) primers. The amplicons of 6 positive colonies per pool were subjected to direct sequencing (24) using the T3 (5' AATTAACCCTCACTAAAGGG) and T7 primers. The DNA sequences obtained were assembled using the GCG package, translated and aligned against the rat VR1 gene using the Blast tools. After analysis, the 3079bp amplicon was shown to have 2 introns of 603bp and 1221bp. The latter intron was located at the 3'end of the PCR product. The coding sequence covered 1255 bp and was separated by the former intron. Therefore the clone EST1 was likely to be a partially spliced and incomplete cDNA.

The clone belonging to cluster 1b (EST3) and derived from a kidney cDNA library was ordered and sequenced using the Bal-31 technique. After assembly of the sequences using the GCG package an identical overlap was identified with the DNA sequence of the cluster hVRc. Moreover a 3'end with a polyadenlyation signal and tail was identified. The complete sequence of the combined hVRb Bal-31 derived sequence and hVRc was 2063 bp (1020 bp of coding and 1043 bp of 3' untranslated sequence).

Reference Example B3: Amplification of the middle section of hVR1 using the Polymerase Chain Reaction

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We formulated the hypothesis that both sequences (hVRa and hVRb/c) were part of a common transcript. If the human and rat VR1 were going to be similar, the 2 contigs should be separated by a gap of approximately 275bp. Primers were designed on both sides of the gap to amplify mRNA from brain tissues in order to clone the gap. A smear was obtained with the sense primer (5' TCTACTTCGGTGAACTGCCC) and antisense (5' ACGGCAGGGAGTCATTCTTC). For specificity 50ng of the PCR product were amplified with the nested sense (5' CTGCAGAACTCCTGGCAGA) and antisense (5' GTCACCACCGCTGTGGAAAA) primers. The 900bp nested amplicon was sequenced and shown to be identical to hVRa at one end and

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hVRb/c at the other end. The middle part of the PCR product was homologous to the rat VR1 sequence. This region corresponded to 91 amino acids. When the sequences of hVRa, hVRb/hVRc and the internal amplicon are combined the total length of the Open Reading Frame (ORF) is 824 amino acids followed by a 3' untranslated sequence of 1043 bp. The human amino acid sequence is 87% identical to the rat sequence over that part of the coding region. This sequence was termed hVR1 because of its high degree of identity with the rat VR1 sequence.

10 Reference Example B4: Isolation of the 5' Terminus of hVR1 by PAC isolation

Since no start codon was identified at the 5' end an additional strategy was designed to identify the full-length sequence. Two primers, sense (5' TCCTCTGGCTTCCAACCCGTT) and antisense (5" GAACTGGGCAGAAAGTGCCT) were designed to amplify a 150bp product from the first intron mentioned in reference example B2. A P1 Artificial Chromosome (PAC) genomic clone (25) was isolated by PCR screening of a PAC library (Genome Systems, St Louis, Missouri). PAC DNA was recovered by using standard plasmid isolation protocol (26). An anti-sense primer was designed (5' CTGGAGTTAGGGTCTCCATCC) to sequence the genomic clone towards the potential 5' end of the gene. An open reading frame with a starting codon was identified. The gene structure was confirmed by using the GenScan software (27). The complete gene has a nucleotide sequence of 2517bp (figure 2) and encoded a 839 amino acid protein (Figures 3 and 4). The gene was named hVR1. Multiple alignment of the amino acid sequence of hVR1 and rat VR1 shows a remarkable degree of identity and similarities between both sequences (figure 5). The rVR1 and hVR1 amino acid sequences are 86% identical. Moreover after protein analysis 6 trans-membrane domains and 3 ankyrin binding domains were identified in hVR1 as in the rat VR1 gene.

Example 1: Full-length Amplification of hVR1 from human DRG and assembly into cloning vectors

HVR1 was PCR amplified in three sections from human DRG template. The 5' fragment was amplified using a sense primer encoding a Notl site and a strong

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Kozak motif followed specific (5) by gene sequence GTCATAGCGGCCGCCGCCACCATGAAGAAATGGAGCAGCAC) and antisense primer (5' AGGCCCACTCGGTGAACTTC). The thermo-cycling conditions used for this amplification included a hot start at 94°C for 4 mins. followed by 35 cycles of 94°C for 1 min, 54°C for 1 min and 72°C for 1 min. A final extension step of 72°C for 5 min completed the reaction. The resulting PCR products were separated on a 2% agarose gel and cloned into pCR®II-TOPO according to the manufacturers instructions supplied with the TOPO™ TA Cloning® kit (Invitrogen). The middle section of hVR1 was PCR amplified using the sense primer: 5' GACGAGCATGTACAATGAGA and antisense primer: 5' GTCACCACCGCTGTGGAAAA. The cycling conditions included a hot start at 94°C for 4 mins, followed by 35 cycles of 1 min at 94°C, 56°C and 72°C. A final extension step of 72°C for 5 min completed the reaction. approximately 870 bp was excised from a 2 % agarose gel and cloned as detailed by the TOPO™ TA Cloning® kit into the vector pCR2.1®-TOPO. Finally the end PCR amplified with the sense primer: was **TGTGGACAGCTACAGTGAGA** and the antisense primer: 5'TGCACTGAATTCGAGCACTGGTGTTCCCTCAG which encoded an EcoRI site for cloning. The PCR conditions included a 90 sec hot start at 94°C followed by 35 cycles of 94°C for 50 sec, 50°C for 50 sec and 72°C for 50 sec. The cycling was completed with a 72°C step for 5 min. PCR products were separated on a 2% agarose gel and cloned into the vector pCR2.1®-TOPO.

Resulting clones for each of the three hVR1-fragments were taken for sequence analysis and separate clones coding a consensus sequence were used in the full length assembly of the gene. The Notl/Dralll (New England Biolabs) digested 5' end fragment ligated together with the middle Dralll/EcoRl fragment into a Notl/EcoRl restricted pBluescript SK (+) vector (Stratagene). Finally, the remaining 3' fragment was introduced into the resulting construct via Mscl and EcoRl restriction sites, a map of the resulting construct is displayed in figure 6A.

Several clones were selected for sequence analysis to confirm that constructs still encoded the hVR1 consensus sequence. These were then digested with Notl/EcoRI and ligated into the mammalian expression vector pCIN5-new (a modified version of pCIN1 (32) having an IVS deletion as well as a 36 bp

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deletion repositioning the start codon of neomycin phosphotransferase immediately after the upstream EMVC IRES) as illustrated in figure 6B.

Example 2: Chromosomal Localisation

The primers used to isolate the PAC clone (reference example B4) were selected for PCR on the G3 radiation hybrid panel from Stanford commercially available from Research Genetics (Huntsville, Alabama). The positive lanes and negative patterns were analysed using the public web server at Stanford University (http://www-sghc.stanford.edu). After analysis the hVR1 gene appears to be located on human chromosome 17 around marker SHGC-36073 (lod score=9.55).

Example 3: mRNA Distribution

The tissue distribution of hVR1 was established by slot-blot hybridisation. RNA was transferred onto a sheet of GeneScreen hybridisation transfer membrane (DUPONT) sandwiched in a slot blotter by suction via a vacuum pump. Once the membrane was rinsed in 2x SSC (3M sodium chloride and 0.3M sodium citrate pH7) for 2 min it was exposed to UV using an Ultraviolet crosslinker (Amersham Life Science) for 1min at 15000uW/cm² thus enabling cross-linkage of the RNA onto the membrane. The amounts of RNA on the blot are unknown. The probe was obtained by PCR amplification of a 260 bp product of the coding region of hVR1 with the following two primers: 5' TGTGGACAGCTACAGTGAGA and 5' GTGGAAAACCCGAACAAGA. Membranes were hybridised for 4 hr shaking at 60°C in a 10% dextran sulphate, 1% SDS (sodium dodecyl sulphate) and 1M NaCl solution. The probe was labelled with [α32P]dCTP (Amersham) using the Rediprime™DNA labelling system (Amersham), so as to obtain approximately 500,000cpm of the labelled probe per ml of prehybridisation solution. Briefly 100ng of probe was boiled for 3 minutes (denaturization) and then cooled on ice for 2 minutes in a total volume of $45\mu l$. This was added to the labelling tube from the kit together with 3µl of 32P dCTP followed by an incubation at 37°C for 30 minutes. 400µl of Herring Sperm DNA (Sigma) at a concentration of 8µg/ml was added to the labelled probe and heated at 99°C for 3 minutes followed by rapid cooling on ice. The labelled probe was added and mixed well in pre-hybridisation solution. The membranes were hybridised overnight at 55°C.

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The membranes were then washed, first at room temperature in 2xSSC and 1% SDS for 5 minutes, followed by 2x SSC and 1% SDS for 30 min at 50°C. If necessary further washes with 1x SSC and 0.5% SDS or 0.1xSSC and 0.1% for 30 mins at the same temperature were carried out. The membranes were then exposed to Scientific Imaging Film AR (Kodak) using intensifying screens at -70° C overnight and the film developed.

The results are shown on figure 7. Strong signals were observed with the positive controls (slots 4B and 5B). Signals are detected on the human DRG slots (1A and 1B). No signals were detected with the water control (slot 3B). Three multi-tissue northern blots (Clontech) with a wide range of tissues have also been hybridised with the same probe, however no signals were detected. RT-PCR was performed on various tissues with the primer combination used to amplify the probe. A strong band was detected in DRG RNA. Taken together these hybridisations suggest that hVR1 is specifically expressed in neuronal tissue and DRG in particular.

Example 4: Design and production of Anti-hVR1 Antibody

The CHIFTTRSRTRLFGKGDSEEASC (peptide68) CGSLKPEDAEVFKDSMVPGEK (peptide69) were synthesised by standard solid phase techniques and purified by gel filtration chromatography. These peptides were conjugated via their Cys residues to the carrier protein, Tuberculin PPD (purified protein derivative) using sulpho-SMCC (sulfosuccinimidyl 4-[Nmaleimidomethyl]-cyclohexan-1-carboxylate). Rabbits, previously sensitised to Bacillus Calmette Guerin (BCG), were inoculated with the resulting conjugates emulsified in incomplete Freund's adjuvant at approx monthly intervals. Serum was prepared from blood samples taken 7 days after each immunisation. The specific antibody response was followed by indirect enzyme-linked immunosorbent assay (ELISA) using free peptide as antigen. Immunoglobulins were purified from high titre sera using immobilsed peptide affinity columns (sulpholink Pierce). Rabbits designated M143, 144 and 145 received peptide68 conjugate, rabbits M146, 147 and 148, peptide69 conjugate.

The antibodies have been validated by specific staining of the recombinant protein expressed in HEK293 cells. Whole cell lysates were prepared in Sample

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Buffer (4 ml dH₂O, 1 ml 0.5 M Tris-HCl, pH 6.8, 0.8 ml glycerol, 1.6 ml 10 % w/v SDS, 0.4 ml 2- β mercaptoethanol and 0.2 ml of 0.05 % w/v bromophenol blue) and proteins separated by SDS-PAGE and transferred to a nitrocellulose filter by electroblotting. Following incubation with the antisera, bound immunoglobulins were revealed using HRP-conjugated secondary antibodies and enhanced chemiluminescence (ECL) detection. The antisera showed specific binding to a protein(s) of the appropriate molecular weight(s) in extracts of VR1 transfected cells, but not in control extracts, this is illustrated in figure 8.

10 Example 5: Insitu localisation of hVR1 using specific antibody

The purified immunoglobulins have been used for immunohistochemical staining of rat DRG tissue sections. Fixed cryosections of DRG were incubated with antibodies for 48h at 4°C at concentrations between 0.1 to 0.5µg/ml. Following a washing step, bound antibodies were detected by indirect immunofluorescence. The antibodies recognised exclusively small diameter cell bodies of the peripheral sensory neurones as displayed in figure 9. This observation has been extended to human DRG tissues for the anti-peptide68 peptide antibodies demonstrating cross-reactivity with the human sequence as expected. Figure 10A demonstrates labelling of DRG cell bodies with an arrow that points to small diameter neuronal cell body) and in figure 10B the arrow points to labelled neurones innervating human skin.

Example 6: Mammalian Cell Expression (examples 6a-6b)

25 Example 6a: Transient expression of hVR1 in mammalian cells

HEK293 cells were plated onto a 6 well plate, containing poly-l-lysine coated coverslips, at 5 x 10^4 cells per well. Next day, fresh media was added to the cells (50% confluent). CalPhos Mammalian Transfection Protocol (Clontech, K2051-1) was used for DNA transfection. For each well of cells, solution A was made up containing 8ug hVR1pCIN5, 2µg pEYFP-N1 reporter DNA, 12.4 µl calcium solution and water to 100μ l. Solution B (hepes buffered saline) was slowly vortexed while solution A was added dropwise. The mixture was incubated at room temperature for 20 minutes, and then added to cells. The plate was slowly rocked to distribute the solution. The cells were incubated at 37° C for 5 hours, and then washed with phosphate buffered saline. Fresh culture

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medium was added and the plate was incubated 24-48 hours for functional analysis.

Example 6b: Stable expression of hVR1 in mammalian cells

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HEK293 cells were plated onto a 6 well plate at 1 x 10⁵ cells per well. Next day, fresh media was added to the cells (50% confluent). CalPhos Mammalian Transfection Protocol (Clontech, K2051-1) was used for DNA transfection. For each well of cells, solution A was made up containing $2\mu g$ hVR1pCIN5, $12.4\mu l$ 2M calcium solution and water to 100µl. Solution B (hepes buffered saline) was slowly vortexed while solution A was added dropwise. The mixture was incubated at room temperature for 20 minutes, and then added to cells. The plate was slowly rocked to distribute the solution. The cells were incubated at 37°C for 5 hours, and then washed with phosphate buffered saline. Fresh culture medium was added and the plate was incubated 48 hours at 37°C, 5% CO2. Cells were harvested into 100mm dishes in selection medium containing 800μg/ml geneticin. Cells were then incubated and fed at 4 day intervals. In total around 10 days selection is required for each single cell to multiply into a visible clone. Well-separated clones were each picked (with a gilson tip) into separate wells of a 96 well plate, containing maintenance medium (400µg/ml geneticin). Cells were expanded into flasks for freezing stocks and functional analysis. Stable cells may be plated at 1 x 10⁵ cells onto poly-I-lysine coated coverslips in 6 well plate, for calcium imaging next day.

Example 7: Functional Analysis of hVR1(examples 7a-7c):

Example 7a: Electrophysiology using patch clamp methods

The activation of human VR-1 channels transiently expressed in HEK293T cells by capsaicin was investigated. Cells grown on poly-L-lysine-coated glass coverslips were placed in a recording chamber (0.5ml) and superfused with extracellular solution (2ml min⁻¹). The extracellular solution contained: NaCl (140mM), KCl (5mM), MgCl2 (2mM), CaCl2 (2mM), 4-(2-hydroxethyl)-1-piperazineethanesulphonic acid (HEPES, 10mM) and glucose (10mM). The pH was adjusted to 7.4 with NaOH and osmolarity ranged from 310-320mOsm l⁻¹. Patch pipettes (borosilicate glass) were pulled using a Sutter P-97 electrode puller. The pipettes were filled with an internal solution consisting of: CsCl

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(140mM), ethylene glycol-bis(β-aminoethyl ether) N,N,N',N'-tetra acetic acid Cs salt (Cs-EGTA, 5mM) and HEPES (10mM). The pH was adjusted to 7.25 using CsOH and the osmolarity ranged from 275-290 mOsm. When filled with this internal solution, patch electrodes had resistances of 2-5 M Ω . Currents were recorded using standard whole-cell voltage clamp recording techniques (31) at room temperature (21-23°C) using an Axopatch 200A amplifier and signals were sampled at 2 or 0.1 kHz. The majority of series resistance errors (80-85%) were minimized with compensation circuitry. Membrane potentials were not corrected for junction potentials (<4 mV). Voltage pulses and data collection were performed on-line using pClamp8 software (Axon Instruments) interfaced with amplifiers. Membrane potentials were maintained at –60mV between protocols.

Capsaicin or capsazepine (CPZ) were applied, using a 'fast-flow sytem', directly onto the recording cell (<1s to equilibrate). The effects of capsaicin were measured either by application during constant recording while holding the membrane potential at -60mV to elicit an inward current, or applying voltage ramps (-100 to +60mV) in the absence and presence of capsaicin. Similarly both these methods of recording currents evoked by the application of capsaicin were used to demonstrate the blockade by the antagonist (CPZ).

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Figure 11A reveals that application of capsaicin (1 μ M), on human VR1 channels transiently expressed in HEK293T cells, produces an inward current when the membrane was held at a potential of -60mV. This response was abolished by 1 μ M CPZ and the blockade was partially reversible.

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In the presence of 1 μ M capsaicin, voltage ramps (-100 to +70mV) produced a current-voltage relationship demonstrating a substantial outward rectification. Addition of 1 μ M CPZ completely blocked the current (figure 11B). Again, only partial recovery was observed, especially for the inward currents evoked by negative potentials.

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Capsaicin-induced desensitisation of human VR-1 channels in the presence of 2mM external calcium is illustrated in figure 12. Voltage ramps (-100 to +70) were applied and the addition of capsaicin (1 μ M) evoked an outwardly rectifying current. Repeated additions of capsaicin resulted in a progressive 'rundown' in

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the size of the response (figure 12A). Figure 12B shows a plot of the current elicited at a potential of +65mV against time illustrating the 'rundown' in current amplitude. Voltage ramps were applied every 20s and capsaicin added at 2min intervals for approximately 40s. By the 6th addition the current had reduced about 4-fold.

When the external calcium was replaced with 5mM EGTA the size of the current increased dramatically (figure 12C). However, when calcium was re-applied to the external solution, the current evoked by capsaicin (1µM) was approximately equivalent to that of the 6th addition shown in (figure 12A).

Example 7b: Calcium Imaging with HEK293 expressing hVR1

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HEK293 cells expressing hVR1 transiently or stably, were plated onto poly-llysine coated cover slips at 1 x 10⁵ cells per well. They were analysed on the following day by calcium imaging (QuantiCell 700, Applied Imaging). On the day of experiment, WASH buffer was prepared by adding CaCl₂ to extracellular medium (ECM) to a final concentration of 2mM, (ECM contains 125mM NaCl, 5mM KCl, 2mM MgCl₂, 0.5mM NaH₂PO₄, 5mM NaHCO₃, 10mM Hepes, 10mM glucose, 0.1% BSA, pH7.4). The calcium sensitive dye solution was prepared by adding 50µl 5% pluronic F-127 in DMSO (Molecular Probes) to a vial of fura2-AM (Molecular Probes). After mixing, 20µl of the fura2-AM solution was added to 10ml WASH. 1.5 ml was then added to cells, which were then incubated at 37°C for 30 minutes. The plate was washed three times with WASH. 1ml WASH was added and stored in dark. Agonists and antagonists were prepared in WASH at 5x their required assay concentrations. The reagents and assay temperature was kept at 37°C. For the transiently transfected cells, the YFP reporter DNA fluorescence (490nm excitation) was used to identify the transfected cells. Cells were initially imaged in 400μl WASH (or 300μl WASH plus 100μl antagonist e.g. capsazepine). After approximately 1 min, 100µl agonist (e.g. capsaicin, anadamide or resiniferatoxin) at 5 x the desired concentration was added to give final 1x concentration. A sequence of images (340/380nm excitation) were taken to monitor calcium influx response in cells before (30-60 secs), and after the addition of agonist (2-5 mins). Figure 13 displays time courses taken for each of the tests set up to look at the affect of the different agonists mentioned above in the presence or absence of the rat VR1 antagonist, capsazepine. The Imager

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also plots graphs of respective calcium concentration (nM) versus time (seconds) as shown in figure 14. After the addition of agonist (e.g. capsaicin, indicated by the vertical arrow on graph), the cells expressing hVR1 are stimulated to influx calcium. This is shown by the appearance of peak on the trace. The peak height correlates with hVR1 expression level. Varying levels of expression is some times seen depending on which cells are selected for the graph. Similar experiments may be accomplished to examine the response of protons and heat.

10 Example 7c: Use of a FLIPR assay with VR1

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FLIPR (Fluorometric Imaging Plate Reader) is a high throughput fluorescencebased drug discovery tool for functional cell analysis. Intracellular calcium is monitored with the calcium sensitive dye, fluo3-AM. HEK293 cells stably expressing rat VR1 were plated into a 96 well, poly-l-lysine treated FLIPR plate at 3 x 104 cells per well. On the following day, the plate was processed for FLIPR. FBP buffer was prepared (15μM Probenecid (calcium ATPase pump blocker) in 1x FLIPR buffer (145mM NaCl, 5mM KCl, 1mM MgCl2, 2mM CaCl2, 10mM glucose, 20mM Hepes). FBP buffer pH was then adjusted to 7.4 with NaOH. 400μl DMSO was added to a vial of fluo3-AM (Cambridge Bioscience, F-1241). The fluo3-AM solution was incubated at 37°C for 10 min and vortexed. LOAD was prepared by adding 20µl of fluo3-AM solution and 20µl 20% pleuronic F-127 in DMSO (Cambridge Bioscience, P-3000) into 10 ml FBP. The 96 well plate containing cells was flicked off to remove cell medium. 100µl LOAD was added per well. Cells were then incubated at 37°C for 60 minutes. Capsaicin (a rVR1 agonist) and capsazepine (CPZ, a rVR1 antagonist) were prepared at 10x the desired final assay concentrations in FBP. The plate was flicked to remove LOAD from cells, and 180µl FBP was added per well. The FLIPR machine added 20µl capsaicin per well to give a final 1x concentration. Cells were monitored for 70 seconds after agonist addition. The FLIPR traces (fluorescence change (counts) versus time (seconds)) were produced for each well. Peaks indicate capsaicin-gated calcium influx, by cells expressing rVR1. The peak height correlates with the rVR1 expression level. To measure antagonism of the VR1 response 20µl 10x antagonist CPZ was added into wells to give a final 1x concentration. The plate was incubated for 15 minutes at room temperature prior reading in the FLIPR. The FLIPR traces recorded for each well show that the

peak heights are reduced in cells pre-incubated in CPZ. The same FLIPR assay may be used to monitor the response of human VR1 on exposure to agonists and antagonists.

5 Example 8: Example of a screen using human VR1.

FLIPR assay technology may be utilised to screen for hVR1 modulators according to the procedure described in figure 15. Human VR1 may be gated with protons, capsaicin or heat.

Reference Example C: Identification and partial characterisation of additional human vanilloid receptors (referenence examples C1-C3):

Reference Example C1: Identification and characterisation of a novel vanilloid-like receptor, hVR3

ESTs belonging to the remaining clusters were characterised by *in silico* cloning (reference example A). The following clones were used during this process: - EST6/EST7 (hVRd), -EST8. (hVRe), - EST9/EST10. (hVRf). These EST clusters have been aligned with rat VR1 in figure 16, note that this diagram is not to scale.

Reference Example C2: Sequencing of clones

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Further sequencing, as detailed in reference example B2, and *in silico* cloning, enabled clusters hVRd, hVRe and hVRf to collapse forming a single contig of 583 amino acids. This sequence was named hVR3 and has 49 % identity with the rat VR1 sequence. It was unlikely that this single contig was a full-length vanilloid receptor transcript as no obvious starting codon was present and it was shorter than the rat VR1 transcript.

Reference Example C3: Identification of the 5' terminus of hVR3

Two primers (sense primer 5' ATGGCCACCAGCAGGGTTAC and antisense primer 5' TCTGCCAGGTTCCAGCTG) designed to PCR amplify an amplicon stretching the 3' end of hVR3 and its 3'utr were used to isolate a genomic PAC clone (Genome Systems. St Louis, Missouri). The hVR3 specific PAC clone was then used as template to generate a library. This was achieved by sonicating 6μg of Qiagen purified PAC construct, size sel cting fragmented DNA of 500-

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2000bp. These resulting fragments were then blunt ended and cloned into the vector pCR[®]-Blunt as detailed in the manufacturers protocol supplied with the Zero Blunt™ PCR cloning kit (Invitrogen). Clones were then sequenced (reference example B2) to identify the complete 5' end of the hVR3 transcript. The full-length nucleotide sequence of the hVR3 gene is displayed in figure 17. Figure 18 illustrates both nucleotide and encoded amino acid sequence of the human VR1 and figure 19 depicts the amino acid sequence of the hVR3 gene with shaded regions denoting predicted trans-membrane regions (boxed) and the ankyrin binding domains (unboxed).

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Example 9: Full-length Amplification of hVR3 from human kidney template

Human kidney was used as a source of template for the PCR amplification of hVR3. Primers used for amplification were designed to isolate the gene in three fragments. Primers designed to isolate the 5' end included a sense primer encoding a Not! site and a strong Kozak motif followed by gene specific sequence (5' GTCATAGCGGCCGCGCGCCACCATGCCCAGGGTAGTTGGAC and antisense primer (5' CACCTCTTGTTGTCACTGGA). The PCR conditions used were a hot start at 94°C for 4 mins, followed by 35 cycles of 94°C for 1 min, 56°C for 1 min and 72°C for 1 min and finally one cycle at 72°C for 5 min. The resulting PCR products were separated on a 2% agarose gel and cloned into pCR®II-TOPO according to the manufacturers instructions supplied with the TOPO™ TA Cloning® kit (Invitrogen). The middle fragment was PCR generated using sense and antisense primers 5' CAAATCTGCGCATGAAGTTCCAG and 5' GCCACGAGAAGTTCCACGTAGTG respectively in the presence of 5% DMSO. PCR thermo-cycling required 35 cycles of 1 min at 94°C, 58°C and 72°C for successful amplification of the fragment which was then excised from a 2% agarose gel for cloning into the pCRII®-TOPO vector. Finally the 3' fragment was amplified with a sense primer 5' GCTGCTCCCATTCTTGCTGA and an antisense primer 5' TGCACTCTCGAGAAATGAGTGGGCAGAGAAGC encoding a Xhol restriction site. This fragment was successfully amplified using a hot start at 94°C for 4 min followed by 35 cycles of 94°C for 50 sec, 48°C for 50 sec and 72°C for 2 min. The cycling was completed with a 72°C step for 5 min. The amplified fragment was excised from a 2% agarose gel and clone into the pCRII®-TOPO vector.

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Resulting clones for each of the three PCR generated hVR3-fragments were taken for sequence analysis and separate clones coding a consensus sequence were used in the full-length assembly of the gene. The DrallI restriction site of the pBluescript SK (+) vector (Stratagene) was firstly abolished by digestion with DrallI followed by a blunt ending step using T₄ DNA polymerase (New England Biolabs). This modified vector was then restricted to enable the ligation of both a NotI/Ncol 5' fragment and Ncol/ EcoRI middle fragment. Finally, the remaining 3' fragment was introduced into the resulting construct via DralII and Xhol sites (figure 20A).

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Several clones were selected for sequence analysis to confirm that the constructs still encoded the hVR3 consensus sequence. These were then digested with Notl/Xhol and ligated into the mammalian expression vector pCDNA3.1 (+) (Invitrogen) as seen in figure 20B. The resulting hVR3 consensus sequence is shown in the multiple alignment along with the full-length sequence of hVR1 and the published hVRL-1 in figure 21.

Example 10: Chromosomal localisation

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The 3' terminus, including the 3' UTR sequence of hVR3 was used to design two primers amplify а product of 360 bp: sense primer 5' ATGGCCACCAGCAGGGTTAC and antisense primer 5' TCTGCCAGGTTCCAGCTG. The G3 radiation hybrid panel from Stanford University (Research Genetics, Huntsville, Alabama) was screened by PCR. The positive and negative lanes were analysed using the public web server at Stanford University (http://www-sghc.stanford.edu). After analysis the hVR3 gene appears to be located on human chromosome 12 around markers D12S177E (lod score=15) and D12S1893 (lod score=14).

Example 11: mRNA distribution

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The following primers (5' ACAAGAAGGCGGACATGCGG and 5' ATCTCGTGGCGGTTCTCAAT) were used to obtain a PCR product from the coding region of hVR3. This amplicon was used as a probe on multi-tissue northern blots, the protocol of which is detailed in example 3, to determine the tissue distribution of the gene (figures 22A, 22B and 22C). A transcript of approximately 3.8 kb was detected in the following tissues (the intensities of the

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signals are indicated in brackets): trachea (very strong), kidney (strong), pancreas (strong), prostate (strong), placenta (strong), bone marrow (weak), adrenal gland (weak), lymph node (weak), spinal cord (weak), thyroid (weak), stomach (weak), lung (weak) and liver (weak).

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Since these commercial blots (Clontech, Palo Alto, California, USA) should have the same amount of RNA it is interesting to note the very strong signal in the trachea lane (figure 22A). This could indicate the potential of hVR3 as a target for respiratory pathologies. It was shown by RT-PCR with the primer combination used to produce the probe that the gene is not expressed in DRG.

Example 12: Riboprobe generation for the in situ localisation of hVR3

The same probe, which was specific to hVR3 in Northern blot analysis (example 11), was used to generate a riboprobe. This hVR3 specific probe was cloned into the T7 and SP6 encoding pCRII®-TOPO vector (Invitrogen). This construct was then used in the *in vitro* transcription of DIG labelled RNA strands from the vectors promoters as described in the manufacturers instructions as detailed in the DIG RNA labelling kit (Roche Molecular Biochemicals). This riboprobe may be used to identify the cellular localisation of hVR3 present in tissues such as trachea, lung, pancreas, prostate, placenta and kidney.

Example 13: Mammalian Cell Expression of hVR3

Expression of hVR3 may be accomplished by transfecting a mammalian cell line such as: HEK283T, HEK293, CHO, COS, HeLa and BHK. A detailed method for both transient and stable transfection is detailed in example 6.

Example 14: Functional Analysis of hVR3

The functional analysis of hVR3 may be studied using the electrophysiology, calcium imaging and FLIPR methods as detailed in examples 7a to 7c.

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Example 15: Example of a drug screen using human VR3.

A stable cell line expressing hVR3 may be used in a drug screen such as a selectivity screen using test compounds that have been identified to have an agonistic or antagonistic action on hVR1. FLIPR assay technology may be utilised to screen for hVR3 modulators as proposed in figure 15.

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Claims

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1. An isolated human vanilloid receptor (hVR) protein or a variant thereof.

- 5 2. An isolated human vanilloid receptor (hVR) protein according to claim 1 which is hVR1 or a variant thereof.
 - 3. An isolated human vanilloid receptor (hVR) protein according to claim 1 which is hVR3 or a variant thereof.
 - 4. An isolated human vanilloid receptor (hVR) protein according to claim 2 having an amino acid sequence as shown in Figure 3.
- 5. An isolated human vanilloid receptor (hVR) protein according to claim 3 having an amino acid sequence as shown in Figure 18.
 - 6. A nucleotide sequence encoding a human vanilloid receptor (hVR) protein or a variant thereof, or a nucleotide sequence which is complementary thereto.
 - 7. A nucleotide sequence according to claim 6 encoding for an hVR1 protein or a variant thereof, or a nucleotide sequence which is complementary thereto.
- 25 8. A nucleotide sequence according to claim 6 encoding for an hVR3 protein or a variant thereof, or a nucleotide sequence which is complementary thereto.
- A nucleotide sequence according to claim 6 which is a cDNA
 sequence.
 - 10. A nucleotide sequence according to claim 7 which is a cDNA sequence
 - 11. A nucleotide sequence according to claim 8 which is a cDNA sequence

- 12. A nucleotide sequence according to claim 7 as shown in Figure 2.
- 13. A nucleotide sequence according to claim 8 as shown in Figure 17.

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- 14. An expression vector comprising a nucleotide sequence according to any one of claims 6 to 13, which is capable of expressing an hVR protein or a variant thereof.
- 10 15. An expression vector according to claim 14 which is capable of expressing an hVR1 protein or a variant thereof.
 - 16. An expression vector according to claim 14 which is capable of expressing an hVR3 protein or a variant thereof.

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- 17. A stable cell line comprising an expression vector according to claim 14.
- 18. A stable cell line comprising an expression vector according to claim 15.
 - 19. A stable cell line comprising an expression vector according to claim 16.
- 25 20. A stable cell line according to claim 17 which is a modified HEK293, CHO, COS, HeLa or BHK cell line.
 - 21. A stable cell line according to claim 18 which is a modified HEK293, CHO, COS, HeLa or BHK cell line.

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- 22. A stable cell line according to claim 19 which is a modified HEK293, CHO, COS, HeLa or BHK cell line.
- 23. An antibody specific for a human vanilloid receptor (hVR) protein or a variant thereof as claimed in any one of claims 1 to 5.

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- 24. An antibody according to claim 23 which is specific for hVR1 or a variant thereof.
- 5 25. An antibody according to claim 23 which is specific for hVR3 or a variant thereof.
 - 26. A method for identification of a compound which exhibits hVR modulating activity comprising contacting a human vanilloid receptor (hVR) protein or a variant thereof according to any one of claims 1 to 5 with a test compound and detecting modulating activity or inactivity.
 - 27. A compound which modulates hVR activity, identifiable by a method according to claim 26.
 - 28. A compound according to claim 27 for use in therapy.
 - 29. The use of a compound according to claim 27 in the manufacture of a medicament for treatment or prophylaxis of a disorder which is responsive to the modulation of hVR activity in a human patient.
 - 30. The use according to claim 28 wherein the disorder is pain, neuropathic pain, inflammatory pain, chronic pain, post-operative pain, rheumatoid arthritic pain, neuropathies, neuralgia, algesia, neurodegeneration, nerve injury, stroke, ischaemia migraine, irritable bowl syndrome (IBS), a respiratory disorder, asthma, chronic obstructive pulmonary disease (COPD), a urological disorder, neuropathy, incontinence, interstitial cystitis or an inflammatory disorder.
- 31. A method of treatment or prophylaxis of a disorder which is responsive to modulation of hVR activity in a human patient which comprises administering to said patient an effective amount of a compound according to claim 27.
- 32. A method according to claim 31 wherein the disorder is pain, neuropathic pain, inflammatory pain, chronic pain, post-operative pain,

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rheumatoid arthritic pain, neuropathies, neuralgia, algesia, neurodegeneration, nerve injury, stroke, ischaemia migraine, irritable bowl syndrome (IBS), a respiratory disorder, asthma, chronic obstructive pulmonary disease (COPD), a urological disorder, neuropathy, incontinence, interstitial cystitis or an inflammatory disorder.

- 33. A compound which modulates hVR activity, identifiable by a method according to claim 26, excluding the compounds capsaicin, resiniferatoxin, piperine, zingerone, polydodial, warburganal, aframodial, cinnamosmolide, cinnamolide, isovelleral, scalaradial, ancistrodial, β-acaridial, scutigeral, merulidial, anandamide and capsazepine.
- 34. A compound according to claim 33 for use in therapy.
- 15 35. The use of a compound according to claim 33 in the manufacture of a medicament for treatment or prophylaxis of a disorder which is responsive to the modulation of hVR activity in a human patient.
- 36. The use according to claim 35 wherein the disorder is pain, neuropathic pain, inflammatory pain, chronic pain, post-operative pain, rheumatoid arthritic pain, neuropathies, neuralgia, algesia, neurodegeneration, nerve injury, stroke, ischaemia migraine, irritable bowl syndrome (IBS), a respiratory disorder, asthma, chronic obstructive pulmonary disease (COPD), a urological disorder, neuropathy, incontinence, interstitial cystitis or an inflammatory disorder.
 - 37. A method of treatment or prophylaxis of a disorder which is responsive to modulation of hVR activity in a human patient which comprises administering to said patient an effective amount of a compound according to claim 33.
 - 38. A method according to claim 37 wherein the disorder is pain, neuropathic pain, inflammatory pain, chronic pain, post-operative pain, rheumatoid arthritic pain, neuropathies, neuralgia, algesia, neurodegeneration, nerve injury, stroke, ischaemia migraine, irritable bowl syndrome (IBS), a respiratory disorder, asthma, chronic obstructive pulmonary disease (COPD), a

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urological disorder, neuropathy, incontinence, interstitial cystitis or an inflammatory disorder.

- 39. A compound identified by the method according to claim 26.
- 40. A compound according to claim 39 for use in therapy.

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- 41. The use of a compound according to claim 39 in the manufacture of a medicament for treatment or prophylaxis of a disorder which is responsive to the modulation of hVR activity in a human patient.
- 42. The use according to claim 41 wherein the disorder is pain, neuropathic pain, inflammatory pain, chronic pain, post-operative pain, rheumatoid arthritic pain, neuropathies, neuralgia, algesia, neurodegeneration, nerve injury, stroke, ischaemia migraine, irritable bowl syndrome (IBS), a respiratory disorder, asthma, chronic obstructive pulmonary disease (COPD), a urological disorder, neuropathy, incontinence, interstitial cystitis or an inflammatory disorder.
- 43. A method of treatment or prophylaxis of a disorder which is responsive to modulation of hVR activity in a human patient which comprises administering to said patient an effective amount of a compound according to claim 39.
- 44. A method according to claim 43 wherein the disorder is pain, neuropathic pain, inflammatory pain, chronic pain, post-operative pain, rheumatoid arthritic pain, neuropathies, neuralgia, algesia, neurodegeneration, nerve injury, stroke, ischaemia migraine, irritable bowl syndrome (IBS), a respiratory disorder, asthma, chronic obstructive pulmonary disease (COPD), a urological disorder, neuropathy, incontinence, interstitial cystitis or an inflammatory disorder.
 - 45. A method of producing an hVR protein or a variant thereof according to any one of claims 1-5 comprising introducing into an appropriate cell line a suitable vector comprising a nucleotide sequence encoding for an hVR protein or

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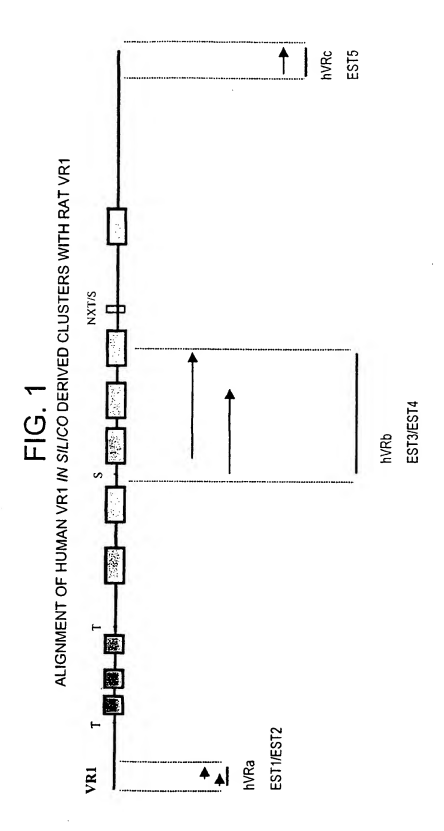
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a variant thereof, under conditions suitable for obtaining expression of the hVR protein or variant thereof.

- 46. A method of producing an hVR1 protein or a variant thereof comprising introducing into an appropriate cell line a suitable vector comprising a nucleotide sequence encoding for an hVR1 protein or a variant thereof, under conditions suitable for obtaining expression of the hVR1 protein or variant thereof.
- 47. A method of producing an hVR3 protein or a variant thereof comprising introducing into an appropriate cell line a suitable vector comprising a nucleotide sequence encoding for an hVR3 protein or a variant thereof, under conditions suitable for obtaining expression of the hVR3 protein or variant thereof.
 - 48. A human vanilloid receptor (hVR) protein or a variant thereof for use in a method of screening for agents useful in the treatment or prophylaxis of a disorder which is responsive to the modulation of hVR activity in a human patient
 - 49. A human vanilloid receptor (hVR) protein according to claim 48 wherein the disorder is pain, neuropathic pain, inflammatory pain, chronic pain, post-operative pain, rheumatoid arthritic pain, neuropathies, neuralgia, algesia, neurodegeneration, nerve injury, stroke, ischaemia migraine, irritable bowl syndrome (IBS), a respiratory disorder, asthma, chronic obstructive pulmonary disease (COPD), a urological disorder, neuropathy, incontinence, interstitial cystitis or an inflammatory disorder.
 - 50. A human vanilloid receptor (hVR) protein according to claim 48 or 49 which is hVR1 or a variant thereof.
- 51. A human vanilloid receptor (hVR) protein according to claim 48 or 49 which is hVR3 or a variant thereof.



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FIG. 2

hVR1 SEQUENCE INCLUDING THE 5'UTR (nt -773 TO nt 0), CODING REGION (nt 1 TO 2517) AND 3'UTR (nt 2518 TO nt 3560)

-773	ccccagccacacacacacacacacacacacacacacaca	-714
-713	aaggccagaagcttgacagatgttgattcataaaaatgcaaaagccaaaatccaaaatct	-654
-653		-594
-593		-534
-533		-474
-473		-414
-413		-354
-353		-294
-293	geegggggeetgteeaccteecaggeegaegteagtggeegeaggaetgeetgggeeet	-234
-233	gctaggcetgctcaggcetctggggtgagaggttcagtcctggaaacacttca	-174
173	gttctagggggctggggcagcagcaagttggagttttggggtaccetgcttcacagggc	-114
113		-54
-53		6
7	AAATGGAGCAGCACAGACTTGGGGGCAGCTGCGGACCCACTCCAAAAGGACACCTGCCCA	66
67	GACCCCTGGATGGAGACCCTAACTCCAGGCCACCTCCAGCCCAGCTCTCCACG	126
127	GCCAAGAGCCGCACCCGGCTCTTTGGGAAGGGTGACTCGGAGGAGGCTTTCCCGGTGGAT	186
187	TGCCCTCACGAGGAGGTGAGCTGGACTCCTGCCCGACCATCACAGTCAGCCCTGTTATC	246
247		306

307	GCCGCCAGCACCGAGAAGACCCTCAGGCTCTATGATCGCAGGAGTATCTTTGAAGCCGTT	366
367		426
427		486
487	ATGCTCAACCTGCACGACGGACAGAACACCACCATCCCCTGCTCCTGGAGATCGCGCGG	546
547		606
607	CAGACAGCACTGCACATCGCCATCGAGAGACGCAACATGGCCCTGGTGACCCTCCTGGTG	666
667	GAGAACGGAGCAGACGTCCAGGCTGCGGCCCATGGGGACTTCTTTAAGAAAACCAAAGGG	726
727	CGGCCTGGATTCTACTTCGGTGAACTGCCCCTGTCCCTGGCCGCGTGCACCAACCA	786
787	GGCATCGTGAAGTTCCTGCAGAACTCCTGGCAGACGGCCGACATCAGCGCCAGGGAC	846
847		906
907		966
967	ACGCTGAAGCTGGAGGAGCTCACCAACAAGAAGGGAATGACGCCGCTGGCTCTGGCAGCT	1026
1027	GGGACCGGGAAGATCGGGGTCTTGGCCTATATTCTCCAGCGGGAGATCCAGGAGCCCGAG	1086
1087	TGCAGGCACCTGTCCAGGAAGTTCACCGAGTGGGCCTACGGGCCCGTGCACTCCTCGCTG	1146
1147		1206
1207	AGCAGCAGCAGACCCCTAATCGCCACGACATGCTCTTGGTGGAGCCGCTGAACCGACTC	1266
1267	CTGCAGGACAAGTGGGACAGATTCGTCAAGCGCATCTTCTACTTCAACTTCCTGGTCTAC	1326
1327	TGCCTGTACATGATCATCTTCACCATGGCTGCCTACTACAGGCCCGTGGATGGCTTGCCT	1386
1387	CCCTTTAAGATGGAAAAAATTGGAGACTATTTCCGAGTTACTGGAGAGATCCTGTCTGT	1446

FIG. 2CONTD

1447 TTAGGAGGAGTCTACTTCTTTTCCGAGGGATTCAGTATTTCCTGCAGAGGCGGCCGTCG 1506 1566 1567 ATGCTGGCCACCGTGGTGCTGTACTTCAGCCACCTCAAGGAGTATGTGGCTTCCATGGTA 1627 TTCTCCCTGGCCTTGGGCTGGACCAACATGCTCTACTACACCCGCGGTTTCCAGCAGATG 1686 GCCATCTATGCCGTCATGATAGAGAAGATGATCCTGAGAGACCTGTGCCGTTTCATGTTT 1746 GTCTACATCGTCTTCTTGTTCGGGTTTTCCACAGCGGTGGTGACGCTGATTGAAGACGGG 1806 1866 1926 1927 ATCGGCATGGGCGACCTGGAGTTCACTGAGAACTATGACTTCAAGGCTGTCTTCATCATC 1986 CTGCTGCTGGCCTATGTAATTCTCACCTACATCCTCCTGCTCAACATGCTCATCGCCCTC 2046 ATGGGTGAGACTGTCAACAAGATCGCACAGGAGAGCAAGAACATCTGGAAGCTGCAGAGA 2106 GCCATCACCATCCTGGACACGGAGAAGAGCTTCCTTAAGTGCATGAGGAAGGCCTTCCGC 2166 TCAGGCAAGCTGCTGCAGGTGGGGTACACACCTGATGGCAAGGACGACTACCGGTGGTGC 2226 TTCAGGGTGGACGAGGTGAACTGGACCACCTGGAACACCAACGTGGGCATCATCAACGAA 2286 GACCCGGGCAACTGTGAGGGCGTCAAGCGCACCCTGAGCTTCTCCCTGCGGTCAAGCAGA GTTTCAGGCAGACACTGGAAGAACTTTGCCCTGGTCCCCCTTTTAAGAGAGGCAAGTGCT 2406 CGAGATAGGCAGTCTGCTCAGCCCGAGGAAGTTTATCTGCGACAGTTTTCAGGGTCTCTG 2466 AAGCCAGAGGACGCTGAGGTCTTCAAGAGTCCTGCCGCTTCCGGGGAGAAGtgaggacgt 2526 2586 2527 cacgcagacagcactgtcaacactgggccttaggagaccccgttgccacgggggggctgct

FIG.2CONT'D

2587	gagggaacaccagtgctctgtcagcagcctggcctggtctgtgcctgcc	2646
2647		2706
2707		2766
2767		2826
2827	ttattcttttctgtgagacagagttcactcttgttgcccaggctggagtgcagtggtgtg	2886
2887	atcttggctcactgcaacctctgctcccgggttcaagcgattcttctgcttcagtctccc	2946
2947	aagtagettggattacaggtgageactaceaegeeeggetaatttttgtattttaatag	3006
3007	agacggggtttcaccatgttggccaggctggtctcgaactcttgacctcaggtgatctgc	3066
3067		3126
3127	tgattttatattattaggagcaaaagtaaatgaagcccaggaaaacacctttgggaacaa	3186
3187	actottectttgatggaaaatgcagaggcccttectetgtgcegtgettgetectett	3246
3247	acctgcccgggtggtttgggggtgtttcctccctggagaagatgggggaggctg	3306
3307		3366
3367		3426
3427		3486
3487		3546
3547	acadatatotatacaaaaaaaaaaaaaaaaaaaaaaaaaa	

FIG. 2contd

FIG. 3

NUCLEOTIDE AND AMINO ACID SEQUENCE OF hVR1 INCLUDING THE 5'UTR (nt -773 TO nt 0), CODING REGION (nt TO 2517) AND 3'UTR (nt 2518 TO nt 3560)

-773	ccccagecacacacacacacacacacacacacacacacac	-714
-713	aaggccagaagcttgacagatgttgattcataaaaatgcaaaagccaaaatccaaaatct	-654
-653	tgtataageteagtggetgtggcagegaggttgaagageaaaggeaggeegggeacetgg	-594
-593	ctgatgatgtgtggacccgttgcacagcagggcccgcagtgcggtgtggggtgtgggg	-534
-533	ccagtctctgccgctcaccctattccagggacacagtctgcttggctcttctggactgag	-474
-473	ccatcctcatcaccgagatcctccctgaattcagcccacgacagccaccccggccgtttt	-414
-413	ccttgttctgtgtgggaagggaggcagcgggtggttatcaacctcaccctgcagaggag	-354
-353	gcacctgaggcccagagacgaggggatgggtctaacccagaaccacagatggctctga	-294
-293	gccgggggcctgtccaccctcccaggccgacgtcagtggccgcaggactgcctgggccct	-234
-233	gctaggcctgctcacctctgaggcctctggggtgagaggttcagtcctggaaacacttca	-174
-173	gttctagggggctgggggcagcaagttggagttttggggtaccctgcttcacagggc	-114
-113	ccttggcaaggaggcaggtggggtctaaggacaagcagtccttactttgggagtcaacc	-54
-53 1	ccggcgtggtggctgctgcaggttgcacactgggccacagaggatccagcaaggATGAAG M K	6 2
7 3	AAATGGAGCAGCACAGACTTGGGGGCAGCTGCGGACCCACTCCAAAAGGACACCTGCCCA K W S S T D L G A A A D P L Q K D T C P	66 22
67	GACCCCCTGGATGGAGACCCTAACTCCAGGCCACCTCCAGCCAAGCCCCAGCTCTCCACG	
23	D P L D G D P N S R P P P A K P Q L S T	126 42
127 43	GCCAAGAGCCGCACCCGGCTCTTTGGGAAGGGTGACTCGGAGGAGGCTTTCCCGGTGGAT A K S R T R L F G K G D S E E A F P V D	186 62
187	TGCCCTCACGAGGAAGGTGAGCTGGACTCCTGCCCGACCATCACAGTCAGCCCTGTTATC	246
63	C P H E E G E L D S C P T I T V S P V I	82
247	ACCATCCAGAGGCCAGGAGACGGCCCCACCGGTGCCAGGGCTGCTGTCCCAGGACTCTGTC	306
83	T I Q R P G D G P T G A R L L S Q D S V	102
307	GCCGCCAGCACCGAGAAGACCCTCAGGCTCTATGATCGCAGGAGTATCTTTGAAGCCGTT	366
103	AASTEKTLRLYDRRSIFEAV	122
367	GCTCAGAATAACTGCCAGGATCTGGAGAGCCTGCTGCTCTTCCTGCAGAAGAGCAAGAAG	426
	AQNNCQDLESLLLFLQKSKK	142
427	CACCTCACAGACAACGAGTTCAAAGACCCTGAGACAGGGAAGACCTGTCTGCTGAAAGCC	486
	H L T D N E F K D P E T G K T C L L K A	162
487	ATGCTCAACCTGCACGACGGACAGAACACCACCATCCCCTGCTCCTCCACATCCCCCC	546

163	M	L	N	L	Н	D	G	Q	N	T	T	I	P	L	L	L	E	I	A	R	182
547	C	ממ.	CCA	CAG	CCT	ממסי	CCA	ССТ	יייבייי	~ ~ ~	CCC	CAG	ста	CAC	CCA	CAC	ርጥ አ	ርጥ አ	~ A A	GGGC	606
183			D			K			V				Y		D		Y	Y	K		202
607	CZ	GAC	AGC	ACT	CA	САТ	יכפס	CAT	CGA	GAG	ACG	CAA	CAT	יככר	сст	יככיי	GAC	CCT	CCT	GGTG	666
203	0	Т	A	L	н	I	A	I	E	R	R		M	A	L	v	Т	L	L	v	222
203	¥	•		-	••	•	-	-		•	• • • • • • • • • • • • • • • • • • • •	••	•••	-	_	•	•		n	•	222
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667																				AGGG	
223	E	N	G	A	ע	٧	Q	A	A	A	н	G	D	F.	F	K	K	T	K	G	242
				·																	
727									_									CAA	CCA	GCTG	786
243	R	P	G	F	Y	F	G	E	L	P	L	S	L	A	A	С	T	N	Q	L	262
787	GC	CAT	CGT	GAA	GTT	CCT	GCI	'GCA	GAA	CTC	CTG	GCA	GAC	GGC	CGA	CAT	CAG	CGC	CAG	GGAC	846
263	G	I	V	ĸ	F	L	L	Q	N	s	W	Q	T	A	D	I	S	A	R	D	282
847	TC	GGT	GGG	CAA	CAC	GGT	GCT	GCA	CGC	CCT	GGT	GGA	GGI	'GGC	CGA	CAA	CAC	GGC	CGA	CAAC	906
283		v							A							N	_		D		302
	_	•	_	••	•	•	_	••	••	_	•	_	•	••	-		•	••	-	••	302
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303		K		v	Т	S	M	Y		E	I	L	I	L	G	A	K	L	H	P	322
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967	20.0	·~~	ר א יי	CCT	יככא	CCN	CCT	יכאכ	~ A A	~ A A	~ N N		אית	יכאכ		·~~m		mem	~~~	AGCT	1026
323	T		K	L	E.	E	L		.CAA N				M M								
	_					_			-					T	P	L	A	L	A	•	342
1027																				CGAG	1086
343	G	T	G	K	I	G	V	L	A	Y	I	L	Q	R	Ε	Ι	Q	E	P	E	362
1087	TO	CAG	GCA	CCT	GTC	CAG	GAA	GTI	CAC	CGA	GTG	GGC	CTA	CGG	GCC	CGT	GCA	CTC	CTC	GCTG	1146
363	С	R	Н	L	S	R	K	F	T	E	W	Α	Y	G	P	v	Н	s	S	L	382
1147	TA	CGA	CCT	GTC	CTG	CAT	CGA	CAC	CTG	CGA	GAA	GAA	CTC	GGT	GCT	'GGA	GGT	GAT	CGC	CTAC	1206
383	Y	D	L	S	С	I	D	T	С	E	K	N	S	V	L	E	V	I	A	Y	402
1207	AC	CAG	CAG	CGA	GAC	ccc	TAA	TCG	CCA	.CGA	CAT	GCT	CTT	'GG'I	'GGA	GCC	GCT	GAA	CCG	ACTC	1266
403	s	S	s	E	T	P	N	R	Н	D	M	L	L	v	E	P	L	N	R	L	422
1267	CI	'GCA	GGA	CAA	GTG	GGA	CAG	ATT	CGT	CAA	GCG	САТ	СТТ	ביים	СТТ	CAA	СТТ	ССТ	GGT	CTAC	1326
423		0							v								F		v		442
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443	c		Y	M	I	I	F	T		A	A	Y	Y	R	P	V	D	G	L	P	462
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1207	~			~~~	~~1					- m s							~		~~~	mama	
1387																					1446
463	Þ	F.	K.	М	E	K	1	G	D	Y	F.	R	V	T	G	E	1	L	S	V	482
1447																				GTCG	1506
483	L	G	G	V	Y	F	F	F	R	G	I	Q	Y	F	L	Q	R	R	P	S	502
1507	AT	GAA	GAC.	CCT	GTT	TGT	GGA	CAG	CTA	CAG	TGA	GAT	GCT	TTT	CTI	TCT	GÇA	GTC	ACT	GTTC	1566
503	M	K	T	L	F	v	D	s	Y	S	E	M	L	F	F	L	Q	s	L	F	522
													_	-	_	_	-	-			
1567	ΑT	GCT	GGC	CAC	CGT	GGT	GCT	GTA	CTT	CAG	CCA	CCT	CAA	GCA	GTA	TCT	GGC	TTC	САТ	GGTA	1626
523																v					542
		_		-	•	•	_	-	-	-		_	- `	-	-	•	••	_		•	
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102/	* *		1	ح ب	CIT	عاوای	~10		~~~		GC I	LIM	CIA	UMU	فاسات	نافات	111	UUM	-	aure	1000

FIG. 3CONTD

543	F S	L	A	L	G	W	T	N	М	L	Y	Y	T	R	G	F	Q	Q	м	562
1687 563	GGCA						AGAT E					GAG R		CCI		CCG R		CAT M	GTTT F	1746 582
	Cmcm:	> C > F			- m	n (** 1777 n	ncci		7 mm/m/	~~~	~ N C (10 x 0	CCM	~ N 17	יוויר א	a (** a	cccc	1806
1747 583	GTCT V Y					F	G		S				v.	T	L	I	E		G	602
1807	AAGA	a mc a	CTC			- C- TP (ייייי	CTI (~~»	~~ m/		\ C \ C					·mcc	-CTC	CACC	1866
603	K N							\$						R						622
1867	cccc		ጥልር	ירייר	ירייי	\	\C \ (عصمة	יחים.	እ ር ጥ/	ירא (רייים	יירריי	יככא	CCT	CTT	ממיי	יייים	CACC	1926
623	P P		S		Y	-			Y				L			F	K		T	642
1927	ATCG	~~ » «		יייי	CCT	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	CTT	יר <i>א</i> נ	-m-	N C N :	N (- 1717)	י שרי	CTTT	ת תיים	ccc	·TOT	سسار	יר א ידי	ር እ ጥ ር	1986
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663	L L													N	-	-				682
2047	N TO C	-m->	C N C	-m	, , ,	· ~ > 1		BCC/	~ > ~ -				~~ ~ 7	C N 17		~ > >	CCT		C	2106
2047 683	ATGG M G		T	V		K			.ΑC Q		AGA S		N		W				R	702
	0001	2010							. ~ .										0000	21.66
2107 703	GCCA'			rcca L			مخاخات E							M M					R	2166 722
																				2225
2167 723	TCAG S G			rgCi L			rgg(G					ATGO G	CA. K		CGA D	CTA Y	CCG R	GTG W	GTGC	2226 742
				_	_															
2227 743	TTCA(GGGI V								CTC W				CGI V		CAI I	CAI I	CAA N		2286 762
743		•	_	_	•	••	**	•	•	••	• • • • • • • • • • • • • • • • • • • •	•	••	•	Ū	-	-	••	-	, 32
2287	GACC																GTC S	AAG S	CAGA R	2346 782
763	D P	G	N	C	E	G	V	K	R	T	L	s	F	s	L	R	3	3	K	762
2347	GTTT																			2406
783	v s	G	R	Н	W	K	N	F	Α	L	V	P	L	L	R	E	A	S	A	802
2407	CGAG	ATAG	GC#	GTC	TGC	TC	AGC	CCG	AGG	AAG!	TTT	ATCI	CCC	ACA	GTI	TTC	AGG	GTC	TCTG	2466
803	R D	R	Q	S	A	Q	P	E	E	V	Y	L	R	Q	F	s	G	S	L	822
2467	AAGC	CAGA	CG2	ACG(TG	\GG1	CT:	CA	AGA	GTC	CTG	CGC	TTC	CGG	GGA	GAA	Gtg	agg	acgt	2526
823	K P	E	D	A	E	V	F	K	s	P	A	A	S	G	E	K				839
2527	cacg	caga	cag	gcac	tgt	caa	acad	etg	ggc	ctt	agga	agad	ccc	egtt	gcc	acç	1999	ggc	tgct	2586
2587	gagg	gaac	acc	agt	gct	cto	gtca	agca	agc	ctg	gcci	tggt	ct	gtgo	ctg	ccc	ago	atg	ttcc	2646
2647	caaa	tctg	tgc	tgç	aca	agg	etgt	-gg	gaa	geg	ttc	ttg	jaaç	gcat	. ggg	gag	jtga	tgt	acat	2706
2707	ccaa																			2766
2767	ctaa	acag	itti	egga	tgg	gtca	agto	ete	tac	tgg	gac	atgt	tag	gged	ctt	gtt	ttc	ttt	gatt	2826
2827	ttat	tctt	ttc	etgt	gaç	gaca	agaç	gtto	cac	tct	tgt	tgco	caç	ggct	gga	gtg	gcaç	jtgg	tgtg	2886
2887	atct	tggc	tca	acto	cae	acct	tet	gcto	ccc	3 99	ttc	agg	gat	tct	tct	gct	tca	gto	tccc	2946

FIG. 3cont'd

2947	aagtagettggattacaggtgagcactaccacgeceggetaatttttgtatttttaatag	3006
3007	agacggggtttcaccatgttggccaggctggtctcgaactcttgacctcaggtgatctgc	3066
3067	ccgccttggcctcccaaagtgctgggattacaggtgtgagccgctgcgctcggccttctt	3126
3127	tgattttatattattaggagcaaaagtaaatgaagcccaggaaaacacctttgggaacaa	3186
3187	actetteetttgatggaaaatgeagaggeeetteetetetgtgeegtgettget	3246
3247	acctgcccgggtggtttgggggtgttggtgtttcctccttggagaagatgggggggg	3306
3307	teccaeteccagetetggcagaateaagetgttgcagcagtgcettetteateett	3366
367	acgatcaatcacagtetecagaagatcageteaattgetgtgcaggttaaaactacagaa	3426
3427	ccacatcccaaaggtacctggtaagaatgtttgaaagatcttccatttctaggaacccca	3486
3487	gteetgetteteegeaatggeacatgetteeacteeatactggeateetcaaataa	3546
3547	acagatatgtatacaaaaaaaaaaaaaaaaaaaaaaaaa	

FIG. 3cont'd

FIG. 4

AMINO ACID SEQUENCE OF hVR1

1	MKKWSSTDLG	AAADPLQKDT	CPDPLDGDPN	SRPPPAKPQL	STAKSRTRLF
51	GKGDSEEAFP	VDCPHEEGEL	DSCPTITVSP	VITIQRPGDG	PTGARLLSQD
101	SVAASTEKTL	RLYDRRSIFE	AVAQNNCQDL	ESLLLFLQKS	KKHL <u>T</u> DNEFK
151	DPETGKTCLL	KAMLNLHDGQ	NTTIPLLLEI	ARQTDSLKEL	VNASYTDSYY
201	KGQTALHIAI	ERRNMALVTL	LVENGADVQA	AAHGDFFKKT	KGRPGFYFGE
251	LPLSLAACTN	QLGIVKFLLQ	NSWQTADISA	RDSVGNTVLH	ALVEVADNTA
301	DNTKFVTSMY	NEILILGAKL	HPTLKLEELT	NHKGMTPLAL	AAGTGKIGVL
351	AYILQREIQE	PECRHLSRKF	T EWAYGPVHS	SLYDLSCIDT	CEKNSVLEVI
401	AYSSSETPNR	HDMLLVEPLN	RLLQDKWDRF	VKRIEYENEL	VYCLYMIÍFT
451	MAAYYRPVDG	LPPFKMEKIG	DYFRVTGEI	SVLGGVYFFF	R GIQY FLQRR
501	PSMKTLFVI S	YSEMLFFLQS,	LEMEATOVLY	es ilkeyvas	MVFSLALGWT
551	NMLYYTRGEQ:	OMGIYAVMI E	KMILRD LCRE	MEVYIVELEG.	ESTAVV <mark>TLIE</mark>
601	DGKNDSLPSE	STSHRWRGPA	CRPPDSSYNS	LYSTCLELFK	FTIGMGDLEF
651	TENYD EKAVF	TITLLLAYVIIT	TYTELLNMET	ALMGETVNKI	AQESKNIWKL
701	QRAITILDTE	KSFLKCMRKA	FRSGKLLQVG	YTPDGKDDYR	WCFRVDEVNW
751	TTWNTNVGII	NEDPGNCXGV	KRTLSFSLRS	SRVSGRHWKN	FALVPLLREA
801	SARDRQSAQP	EEVYLRQFSG	SLKPEDAEVF	KSPAASGEK*	

Key

T/S predicted phosphorylation sites

}: -4 .	Transmembrane domains
	Ankyrin binding domains

FIG. 5

COMPARISON OF THE AMINO ACID SEQUENCE OF THE RAT (VR1) AND HUMAN (hVR1) VANILLOID PROTEINS.

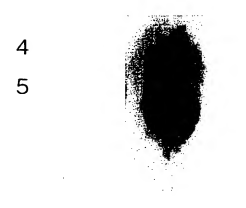
	10	20	30	40	50
VR1	MEQRASLDSEESES				
hVR1	MKKWSSTDLGAAAD				
	60	70	80	90	100
VR1	GKGDSEEASPLDCP				
hVR1	GKGDSEEAFPVDCP		ITVSPVITI	QRPGDGPTGAI	RLLSQD
1770.1	110	120	130	140	150
VR1	SVSAG.EKPPRLYD				
hVR1	SVAASTEKTLRLYD	KRŞIFEAVAQN 170			
VR1	DPETGRTCLLKAMI		180 T.T.T.DVAPRT	190	200 Vmpevv
hVR1	DPETGKTCLLKAML	AND	المرافق وراب	Contract Section 1	
1141/1	210	220	230	240	250
VR1	KGQTALHIAIERRN	MTLVTLLVENG	ADVQAAANG	DFFKKTKGRP	
hVR1	KGQTALHIAIERRN	MALVILLVENG	ADVQAAAHG I	DFFKKTKGRP	GFYFGE
	260	270	280	290	300
VR1	LPLSLAACTNOLAI				
hVR1	LPLSLAACTNOLGI				
VR1	310 DNTKFVTSMYNEIL	320	330 TERTUNDEC	340	350
hVR1	DNTKFVTSMYNEIL	12 A 1 A 1 A 1 A 1 A 1 A 1 A 1 A 1 A 1 A	1 44210	Secretary and a second secretary and the second	
nvki	360	370	380	390	400
VR1	AYILQREIHEPECR				
hVR1	AYILOREIQEPECR	ACCOUNTS TO THE REAL PROPERTY OF THE PARTY O	- 11 CH	and the second of the first terms of the second	4.00 24.5.5.4
	410	420	430	440	450
VR1	AYSSSETPNRHDML				
hVR1	AYSSSETPNRHDML				
VR1	460 AAAYYRPVEGLPPY	470	480	490	500
	MAAYYRPVDGLPPF	CAP. 126 and an array of the second	the second secon	Contraction Contract	
hVR1	510	520	VIGEILSVL(5GVIFFFRGIÇ 540	550 550
VR1	RPSLKSLFVDSYSE		VSVVLÝFSOI	REYVASMVFS	LAMGW
hVR1	RPSMKTLFVDSYSE				
	560	570	580	590	600
VR1	TNMLYYTRGFQQMG	57555 ST \$255 C C C C C C C C C C C C C C C C C C	一、大学、大学、大学、大学、大学、大学、大学、大学、大学、大学、大学、大学、大学、	to the second was to the first the second of	and the same and the same
hVR1	TNMLYYTRGFQQMG		RDLCRFMFV	<i>CIVELEGESTA</i>	VVTLI
VR1	610 EDGKNNSLPMESTP	620	630	640	650
	the state of the state of the state of the state of	r all report from the second	 The rest of the contract of the c	A STANKA AND ANY CAME AND A SECOND	
hVR1	EDGKNDSLPSESTS				
VR1	660 FTENYDFKAVFIIL	670 [.T.A.V.V.TV.T	680 LTNMLTATNA	690 Emilian 1800	700
	FTENYDFKAVFIIL				
HAKT	710	720	730	740	750
VR1	LORALTILDTEKSF				VDEVN
hVR1	LORAITILDTEKSF				
	760	770	780	790	800
VR1	WITWNTNVGIINED	25 25 7 . 1 . A	and the first the commence of	the same of the second section in the second section in the second section is a second section of the second section section is a second section of the second section	
hVR1	WITWNTNVGIINED			GRHWKNFALV	PLLRE
VR1	810 ASTRORHATOOEEV	820	830	aterana:	
	内ではない こうしょうじょう かいしょうしゅん はんかん カル・フェース かんかん	the state of the s	The second secon	CATOM BAZ DOLA !	
hVR1	ASARDRQSAQPEEV	THKALZĞZİKLI	LUALVEKSPA	ASGEK.	

6B -SacII BstEll hVR1 2000 IRES SgrAl /Bsu36 Bbel Ehel Kasi Nari , 000 100 3000 FULL-LENGTH hRV1 CLONED INTO (A) pBLUESCRIPT SK(+) (hVR1pBSK) AND (B) pCIN5-NEW (hVR1pCIN5) VIA Notl/EcoRI RESTRICTION SITES. Notl SnaBl BamHi Ndel i CMV promoter Neo 7413 bps , 2000 4000 V Amp 6000 2000 Miul Bst1107 Muni BgIII. Sspl Scal. Pvul-Ahdl-BC 등 Ball hVR1 Bsu36 1000 Bpull ECORI HindIII Clai Sall Asp718 2000 hVR1pBSK 5470 bps Set 3000 5000 Nacl BsaAI' NgoMi BspLUII 4000 Affill Amp Scal Ahdl ... eA

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FIG. 7
SLOT HYBRIDISATION WITH hVR1 PROBE





Well

1A hDRG 2A rDRG 1B hDRG

3A Water

4B EST3 clone

5B 260bp Amplicon from Brain cDNA

FIG. 8
WESTERN BLOT PROBED WITH ANTI-hVR1 ANTIBODIES.
ARROW POINTS TO hVR1 SPECIFIC BAND

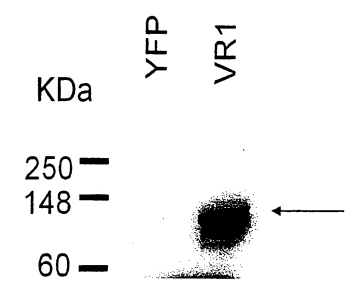


FIG. 9
IN SITU LOCALISATION OF VR1 IN RAT DRG TISSUE SECTIONS.
ARROW POINTS TO A VR1 EXPRESSING SMALL DIAMETER
(<25µn) NEURONE CELL BODY, MAGNIFICATION USED 147x10.

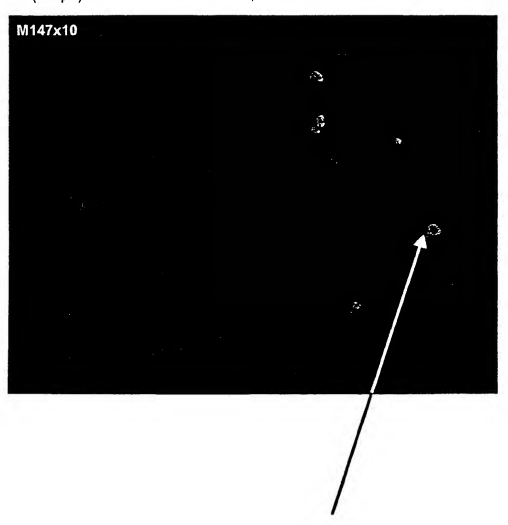


FIG. 10A





FIG. 10B



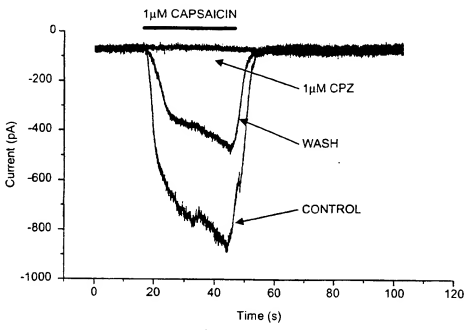
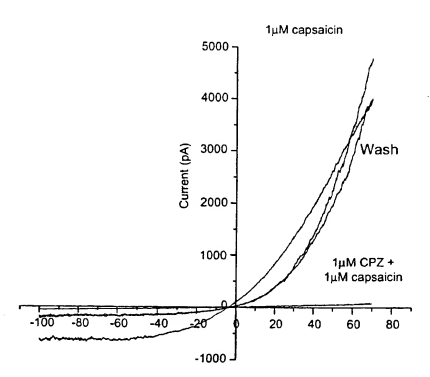
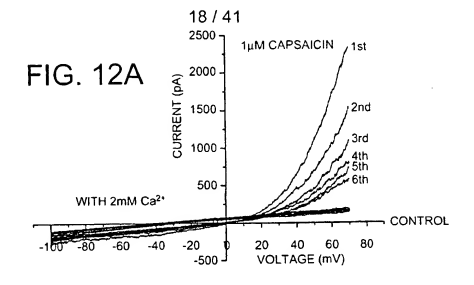


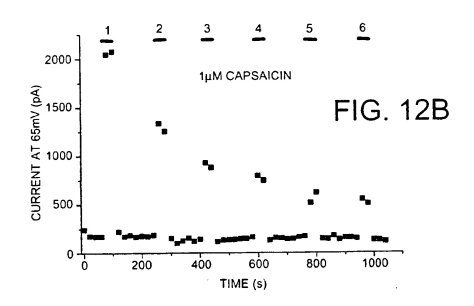
FIG. 11A

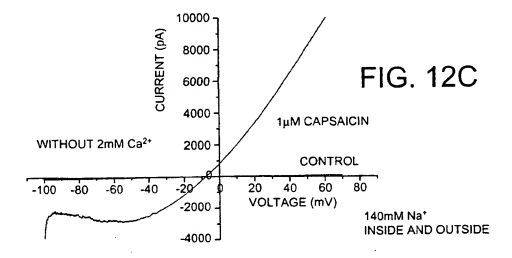


SOLUTIONS OUTSIDE 140mM Na⁺ 2mM Ca²⁺ INSIDE 140mM Cs⁺

FIG. 11B







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13A pCIN5-new in HEK293T, 24hr transient expression, stimulated with 3 μ M capsaicin at time point 52 secs of time course



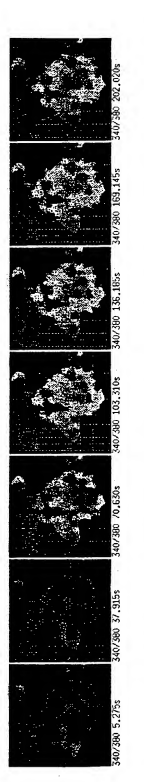
138 hVR1pCIN5 in HEK293T, 24hr expression, stimulated with 1µM capsaicin at time point 52 seconds



13C hVR1pCIN5 in HEK293T, 24hr transient expression, 20 min pre-incubation with 10 uM capsazepine, stimulated with 1 μ M capsaicin at time point 52 seconds of time course



13D hVR1pCIN5 in HEK293T, 24hr transient expression, stimulated with 10uM anandamide at time point 52 seconds



13E hVR1pCIN5 in HEK293T, 24hr transient expression, 20 min pre-incubation in 10uM capsazepine, stimulated with 10uM anandamide at time point 52 sec

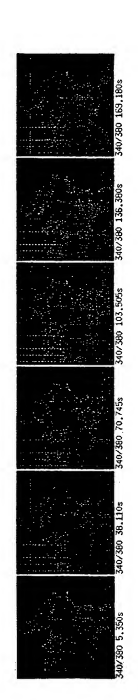
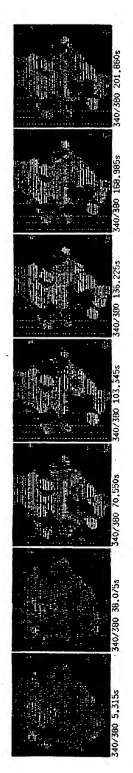


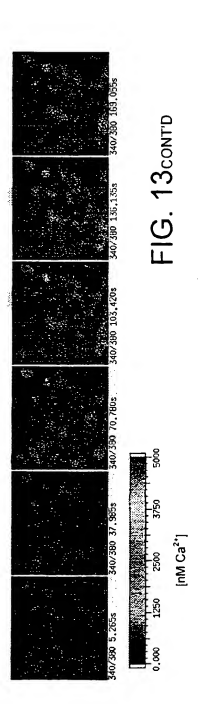
FIG. 13contd



13F hVRIpCIN5 in HEK293T cells, 24hr transient expression, stimulated with 1uM Resiniferatoxin at time point 52 seconds



136 hVR1pCIN5 in HEK293T, 24hr transient expression, 20 min pre-incubation with 10 uM capsazepine, stimulated with 1 uM Resiniferatoxin at time point 52 seconds



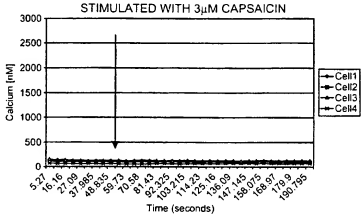
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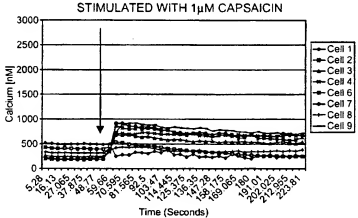
FIG. 14

EXPOSURE OF TRANSFECTED CELLS TO AGONISTS (ADDITION INDICATED BY ARROW).

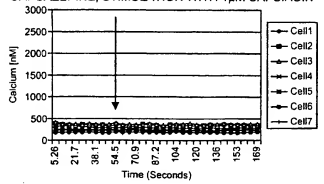
14A: pCIN5-NEW IN HEK293T, 24hr TRANSIENT EXPRESSION,



14B: hVR1pCiN5 IN HEK293T, 24hr EXPRESSION,



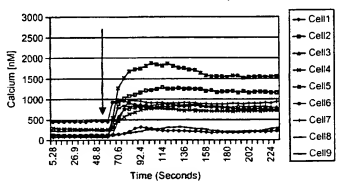
14C: hVR1pCIN5 IN HEK293T, 24hr TRANSIENT EXPRESSION, 20 MIN PRE-INCUBATION WITH 10µM CAPSAZEPINE, STIMULATION WITH 1µM CAPSIACIN



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FIG. 14contid

14D: hVR1pCiN5 IN HEK293T, 24hR TRANSIENT EXPRESSION, STIMULATION WITH 10µM ANANDAMIDE



14E: hVR1pCIN5 IN HEK293T, 24hr TRANSIENT EXPRESSION, 20 MIN PRE-INCUBATION IN 10µM CAPAZEPINE, STIMULATED WITH 10µM ANANDAMIDE

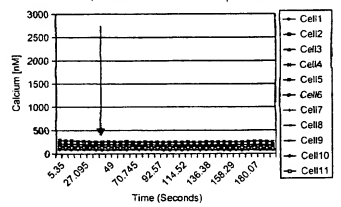
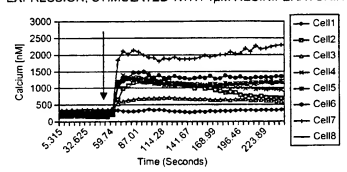
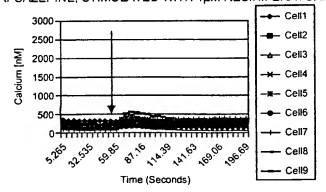


FIG. 14CONTD

14F: hVR1pCIN5 IN HEK293T CELLS, 24hr TRANSIENT EXPRESSION, STIMULATED WITH $1\mu M$ RESINIFERATOXIN



14G: hVR1pCIN5 IN HEK293T, 24hr TRANSIENT EXPRESSION, 20 MIN PRE-INCUBATION WITH 10μM CAPSAZEPINE, STIMULATED WITH 1μM RESINIFERATOXIN



hVR1 ASSAY

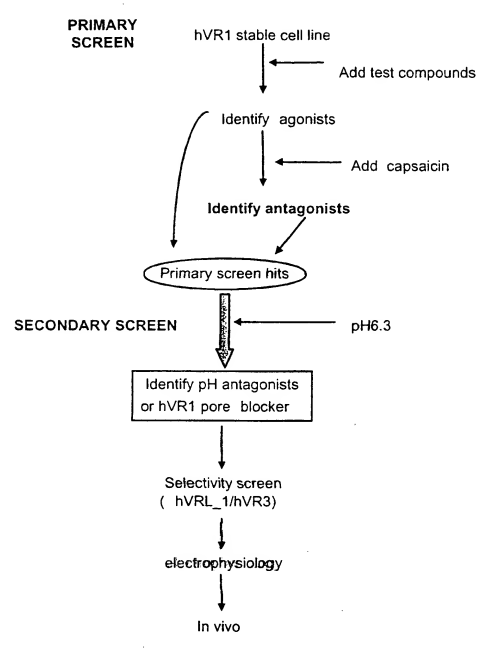
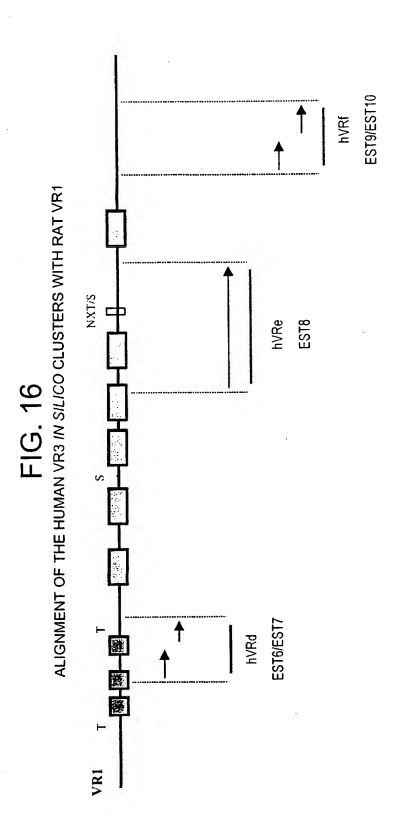


FIG. 15



SUBSTITUTE SHEET (RULE 26)

FIG. 17

hVR3 SEQUENCE INCLUDING 5' UTR (nt -686 TO nt 0) CODING REGION (nt1 TO nt 2889), 3'UTR (nt 2890 TO nt 3418)

-684	ttacgcgttaagaaatacccaagcttatgcatcaagcttggtaccgagctcggatccact	-625
-624	agtacegeeggeeagtgtgetggaattcaaggtgaggaggaggageatggateetgggage	-565
-564	gagtgtgtgcaggccagggaggctttccagaggagcccagttgagctggaacaccagtg	-505
504	gggaggagttgaccagcaaaggtgcagggagggatcagcactttgcactggggagcagag	-445
444	tttgtgcactggggaagtcaactcaagtattggagcctcagtttcctgttctgtaaaatg	-385
384	ggttcatcatgacagtgtttgatgaggaaaaggactgccggcctacacagcaagtccaca	-325
324	tggattttctgagcccctcctgtgcctgaagcccacggttaatggttctgccttagcagg	-265
264	tgcttaccacgtgccaggcactgcactgcactggccactggactgcatgttctgtccatg	-205
204	aggettggatateeceatettacagateaggaagetgaggetatgaaatgtegaettget	-145
144	caatgtcatggaatgactaagtgtggagcctggatttgaacttggctctctggggctcca	-85
-84	aagctggetttettggteageagtagggtetgggateeaagtatggggteecagettgae	-25
-24	cctgaagtccaccctctttcagctaATGCCCAGGGTAGTTGGACCTGGGGCCAATTTGTG	35
36	TTTCCAGGTTCGTGAAAGAGGCTCCTGTTGCAGTTCCCGCCTGAGGCTGGCGGCCAACCA	95
96	CATCTGGGAGTGGCCTCCCTGTGCCCCTGTCATTACAACGGTGGCTTTGAAGCAGCTGGC	155
156	AGCACTGCTGCTGCCACGTGGGAGGGGGCTTCCTGGAGCCCCCGCCCCTGGCCGGGTT	215
216	CTGCCTGACTCCCCTTTCATTCCCTTGCAGGCTGAGCAGTGCAGACGGGCCTGGGGCAGG	275
276	CATGCCGGATTCCAGCGAAGGCCCCCGCGCGGGGGCCCGGGGAGGTGGCTGAGCTCCCCGG	335
336	CCATCACACCACCCACCCACCCTACCCCTACCCCCACCCCCC	205

FIG. 17 CONT'D

1536 CCTCTCCTCCTGGACACGTGTGGGGAAGAGGCCTCCGTGCTGGAGATCCTGGTGTACAA 1595 1596 CAGCAAGATTGAGAACCGCCACGAGATGCTGGCTGTGGAGCCCATCAATGAACTGCTGCG 1656 GGACAAGTGGCGGAAGTTCGGGGCCGTCTCCTTCTACATCAACGTGGTCTCCTACCTGTG 1716 TGCCATGGTTATCTTCACTCTCACCGCCTACTACCAGCCGCTGGAGGGCACACCGCCGTA 1836 TGGGGTCCTGTTCTTCTCACCAACATCAAAGACTTGTTCATGAAGAAAATGCCCTGGAGT 1896 GAATTCTCTCTTCATTGATGGCTCCTTCCAGCTGCTCTACTTCATCTACTCTGTCCTGGT 1956 GATCGTCTCAGCAGCCCTCTACCTGGCAGGGATCGAGGCCTACCTGGCCATGATGGTCTT 2016 TGCCCTGGTCCTGGGCTGGATGAATGCCCTTTACTTCACCCGTGGGCTGAAGCTGACGGG 2076 GACCTATAGCATCATGATCCAGAAGATTCTCTTCAAGGACCTTTTCCGATTCCTGCTCGT 2196 CAACATGAAGGTGTGCAATGAGGACCAGACCGACTGCACAGTGCCCACTTACCCCTCGTG 2256 CCGTGACAGCGAGACCTTCAGCACCTTCCTCCTGGACCTGTTTAAGCTGACCATCGGCAT 2315 2316 GGGCGACCTGGAGATGCTGAGCAGCACCAAGTACCCCGTGGTCTTCATCATCCTGCTGGT 2376 GACCTACATCATCCTCACCTCTGTGCTGCTCCTCAACATGCTCATTGCCCTCATGGGCGA 2436 GACAGTGGCCCAGGTCTCCAAGGAGCAAGCACATCTGGAAGCTGCAGTGGGCCACCAC 2496 CATCCTGGACATTGAGCGCTCCTTCCCCGTATTCCTGAGGAAGGCCTTCCGCTCTGGGGA 2556 GATGGTCACCGTGGGCAAGAGCTCGGACGGCACTCCTGACCGCAGGTGGTGCTTCAGGGT 2616 GGATGAGGTGAACTGGTCTCACTGGAACCAGAACTTGGGCATCATCAACGAGGACCCGGG 2675

FIG. 17 CONTD

2676	CAAGAATGAGACCTACCAGTATTATGGCTTCTCGCATACCGTGGGCCGCCTCCGCAGGGA	2735
2736	TCGCTGGTCCTCGGTGGTACCCCGCGTGGTGGAACTGAACAAGAACTCGAACCCGGACGA	2795
2796	GGTGGTGCCTCTGGACAGCATGGGGAACCCCCGCTGCGATGGCCACCAGCAGGGTTA	2855
2856		2915
2916	ctctgcccactcatttctagtccagccgcatttcagcagtgccttctggggtgtccccc	2975
2976	acaccetgetttggccccagaggegagggaccagtggaggtgccagggaggccccaggac	3035
3036	cctgtggtcccctggctctgcctccccaccctggggtgggggctcccggccacctgtctt	3095
3096	gctcctatggagtcacataagccaacgccagagcccctccacctcaggccccagccctg	3155
3156	cetetecattatttatttgetetgeteteaggaagegaegtgaeceetgeeceagetgga	3215
3216	acctggcagaggccttaggaccccgttccaagtgcactgcccggccaagccccagcctca	3275
3276	geetgegeetgagetgeatgegeeaccatttttggeagegtggeagetttgeaagggget	3335
3336	ggggccctcggcgtggggccatgccttctgtgtgttctgtagtgtctgggatttgccggt	3395
396	gctcaataaatgtttattcattgaaaaaaaaaaaaaaa 3433	

FIG. 17_{CONT'D}

FIG. 18

NUCLEOTIDE AND AMINO ACID SEQUENCE OF hVR3 INCLUDING THE 5'UTR (nt -684 TO nt 0), CODING REGION (nt1 TO 2889) AND 3'UTR (nt 2890 TO nt 3418)

-684	tta	acgo	egtt	caaç	gaaa	ata	ccc	aag	ctta	atgo	ato	caa	gcti	tgg	tac	egaç	gcto	egga	atco	cact	-625
-624	agt	taco	gco	ggg	caç	gtg	tgc	tgg	aati	caa	aggt	tga	gga	gag	gago	cat	ggat	taat	ggg	gage	-565
-564	gag	gtgt	gtg	gcaç	geo	cag	gga	3 99	c t tt	cca	gaç	gga	gaa	cag	ttga	agct	tgga	aca	cca	agtg	-505
-504	9 99	gago	gagt	tga	acca	agca	aaa	ggti	gcag	gga	ggg	gate	cago	cac	ttt	gcad	ctg	gga	ıgca	agag	-445
-444	ttt	gtç	gcac	tgg	gga	aagt	tca	acto	caaç	gtat	tgç	gago	ccto	cagi	ttto	ecto	jtto	etgt	aaa	atg	-385
-384	ggt	tca	tca	tga	cac	j tgi	ttt	gate	gago	gaaa	ago	gacı	tged	egge	ecta	acad	cago	caaç	tcc	aca	-325
-324	tgg	att	ttc	tga	ged	cct	taci	tgt	gcct	gaa	gco	ccad	eggt	ttaa	atgo	jtto	etgo	ctt	ago	agg	-265
-264	tgo	tta	cca	cgt	gec	agg	gca	etgo	cact	.gca	cto	ggc	cact	tgga	acto	gcat	gtt	ctg	rtco	atg	-205
-204	agg	ctt	.gga	tat	ccc	cat	cti	taca	agat	caç	gaa	iget	tgaç	ggct	ato	aaa	tgt	cga	ctt	gct	-145
-144																				.cca	-85
-84	aag	rctg	gct	ttc	ttg	gto	ago	agt	ago	gto	tgg	ga t	cca	agt	ato	1999	tcc	cag	ctt	gac	-25
-24 1	cct	.gaa	gtc	cac	ect	ctt	tea	igct	AA: M		CAG R	GGT V	ragi V	TGC G	P	TGG G	GGC A	CAA N	TTT L	GTG C	35 12
36	TTT	'CCA	GGT	TCG	TGA	AAC	AGO	CTC	CTC	TTG	CAG	TTC	ccc	cci	GAG	GCI	GGC	GGC	CAA	CCA	95
13	F	Q	v	R	E	R	G	s	С	С	s	s	R	L	R	L	A	A	N	Н	32
96	CAT	CTG	GGA	GTG	GCC	TCC	CTC	TGC	ccc	TGI	CAI	TAC	AAC	GGI	'GGC	TTT	'GAA	GCA	GCT	GGC	155
33	I	W	E	W	P	P	С	A	P		I	T	T	v	A	L	K	Q	L	A	52
156	AGC	ACT	GCT	GCT	TGT	CCA	CGI	'GGG	AGG	GGG	CTT	CCI	GGA	GCC	ccc	GCC	CCT	GGC	CGG	GTT	215
53	A	L	L	L	V	H	V	G	G	G	F	L	E	P	P	P	L	A	G	F	72
216	CTG	CCT	GAC	TCC	CCT	TTC	TA	'ccc	TTG	CAG	GCT	GAG	CAG	TGC	AGA	.CGG	GCC	TGG	GGC	AGG	275
73		L	T	P	L	s	F	P			L	S	s	A	D	G	P		A	G	92
276	CAT	GGC	GGA	TTC	CAG	CGA	AGG	CCC	CCG	CGC	GGG	GCC	CGG	GGA	GGT	GGC	TGA	GCT	ccc	CGG	335
93	M			S	s	E	G	₽	R	A	G	P	G	E	V	A	E	L	P	G	112
336	GGA	TGA	GAG	TGG	CAC	CCC	AGG	TGG	GGA	.GGC	TTT	TCC	TCT	CTC	CTC	CCT	GGC	CAA	TCT	GTT	395
113	D	E	s	G	T	P	G	G	E	A	F	P	L	s	S	L	A	N	L	F	132
396	TGA	GGG	GGA	GGA	TGG	CTC	CCT	TTC	GCC	CTC	ACC	GGC	TGA	TGC	CAG	TCG	ccc	TGC	TGG	ccc	455
133	E	G	E	D	G	S	L	S	P	S	P	A	D	A	S	R	P	A	G	P	152
456	AGG	CGA!	TGG	GCG	ACC	AAA	TCT	GCG	САТ	GAA	GTT	CCA	GGG	CGC	ርፓጥ	CCC	CAA	GGG	CCT	GCC	515
153	G	D	G	R	P	N	L	R	M	ĸ	F	Q	G	A	F	R	K	G	V	P	172
516	CAA	CCC	CAT	CGA'	TCT	GCT	GGA	GTC	CAC	CCT	ATA	TGA	GTC	CTC	೧೭೯	೧೭ಌ	حدد	ጥርር:	בררי	CAA	575
173	N	P	I	D	L	L,	E	s	T	L	Y	E	s	s	v	V	P	G	P	K	192

576	GAAAGCACCCATGGACTCACTGTTTGACTACGGCACCTATCGTCACCACTCCAGTGACAA	635
193	K A P M D S L F D Y G T Y R H H S S D N	212
636	CAAGAGGTGGAGGAAGAAGATCATAGAGAAGCAGCCGCAGAGCCCCCAAAGCCCCTGCCCC	695
213	K R W R K K I I E K Q P Q S P K A P A P	232
696	TCAGCCGCCCCCATCCTCAAAGTCTTCAACCGGCCTATCCTCTTTGACATCGTGTCCCG	755
233	Q P P P I L K V F N R P I L F D I V S R	252
756	GGGCTCCACTGCTGACCTGGACGGGCTGCTCCCATTCTTGCTGACCCACAAGAAACGCCT	815
253	GSTADLDGLLPFLLTHKKRL	272
21.6	AACTGATGAGGAGTTTCGAGAGCCATCTACGGGGAAGACCTGCCTG	875
	T D E E F R E P S T G K T C L P K A L L	292
273	T D E E F R E P S I G R I C E F R R E E	232
876	GAACCTGAGCAATGGCCGCAACGACACCATCCCTGTGCTGCTGGACATCGCGGAGCGCAC	935
	N L S N G R N D T I P V L L D I A E R T	312
233		
936	CGGCAACATGCGGGAGTTCATTAACTCGCCCTTCCGTGACATCTACTATCGAGGTCAGAC	995
313	G N M R E F I N S P F R D I Y Y R G Q T	332
996	AGCCCTGCACATCGCCATTGAGCGTCGCTGCAAACACTACGTGGAACTTCTCGTGGCCCA	1055
333	ALHIAIERRCKHYVELLVAQ	352
	GGGAGCTGATGTCCACGCCCAGGCCCGTGGGCGCTTCTTCCAGCCCAAGGATGAGGGGGG	1115
353	G A D V H A Q A R G R F F Q P K D E G G	372
	CTACTTCTACTTTGGGGAGCTGCCCCTGTCGCTGCCTGCACCAACCA	1175
373	Y F Y F G E L P L S L A A C T N Q P H I	392
		1235
1176 393	TGTCAACTACCTGACGGAGAACCCCCACAAGAAGGCGGACATGCGGGCGCCAGGACTCGCG V N Y L T E N P H K K A D M R R Q D S R	412
393	V N I L I E N P N K K A D M K K Q D S K	412
1236	AGGCAACACAGTGCTGCATGCGCTGGTGGCCATTGCTGACAACACCCCGTGAGAACACCCAA	1295
	G N T V L H A L V A I A D N T R E N T K	432
410		
1296	GTTTGTTACCAAGATGTACGACCTGCTGCTGCTCAAGTGTGCCCGCCTCTTCCCCGACAG	1355
	F V T K M Y D L L L K C A R L F P D S	452
1356	CAACCTGGAGGCCGTGCTCAACAACGACGGCCTCTCGCCCCTCATGATGGCTGCCAAGAC	1415
453	N L E A V L N N D G L S P, L M M A A K T	472
	GGGCAAGATTGGGATCTTTCAGCACATCATCCGGCGGGAGGTGACGGATGAGGACACACG	1475
473	G K I G I F Q H I I R R E V T D E D T R	492
	GCACCTGTCCCGCAAGTCCAAGGACTGGGCCTATGGGCCAGTGTATTCCTCGCTTTATGA	
493	H L S R K S K D W A Y G P V Y S S L Y D	512
1526	CCTCTCCCTGGACACGTGTGGGGAAGAGGCCTCCGTGCTGGAGATCCTGGTGTACAA	1595
	L S S L D T C G E E A S V L E I L V Y N	532
513	T 2 2 T T T C G T W 2 A T T T A I W	JJ2
1506	CAGCAAGATTGAGAACCGCCACGAGATGCTGGCTGTGGAGCCCATCAATGAACTGCTGCG	1655
	S K I E N R H E M L A V E P I N E L L R	
		-
1656	GGACAAGTGGCGGAAGTTCGGGGCCGTCTCCTTCTACATCAACGTGGTCTCCTACCTGTG	1715
	D K W R K F G A V S F Y I N V V S Y L C	572

FIG. 18contd

716	TGC																				1775
573									A										P		592
776	CCC	TTA	CCG	CAC	CAC	GGT	GGA	CTA	CCT	GCG	GCT	GGC	TGG	CGA	GGT	CAT	TAC	GCT	CTT	CAC	1835
593	P	Y	R	T	T	v	D	Y	L	R	L	A	G	E	V	I	T	L	F	T	612
836	TGG	GGT	CCT	GTT	CTT	CTT	CAC	CÀA	CAT	CAA	AGA	CTT	GTT	CAT	GAA	GAA	ATG	ccc	TGG	AGT	1895
613	G	V	L	F	F	F	T	N	I	K	D	L	F	M	K	K	С	P	G	V	632
1896	GAA	ттс	TCT	CTT	CAT	TGA	TGG	CTC	CTT	CCA	GCT	GCT	CTA	CTT	CAT	CTA	CTC	TGT	CCT	GGT	1955
633	N	s	L	F	I	D	G	s	F	Q	L	L	Y	F	I	Y	s	V	L	V	652
1956	GAT	CGT	СТС	AGC	AGC	CCT	CTA	CCT	'GGC	AGG	GAT	CGA	GGC	CTA	CCT	GGC	CAT	GAT	GGT	CTT	2015
653	I	v	s	A	A	L	Y	L	A	G	I	E	A	Y	L	A	M	M	V	F	672
2016	TGC	CCT	GGT	CCT	GGG	CTG	GAT	GAA	TGC	CCT	TTA	CTT	CAC	CCG	TGG	GCT	GAA	GCT	GAC	GGG	2075
673	A	L	V	L	G	W	M	N	A	L	Y	F	T	R	G	L	К	L	T	G	692
2076	GAC	CTA	TAG	CAT	CAT	GAT	CCA	GAA.	GAI	TCT	CTI	CAA	GGA	CCI	TTT	CCG	TTA	CCI	GCT	CGT	2135
693																			L		712
2136	CTA	CTI	GCI	CTT	CAT	GAT	CGG	CTA	CGC	TTC	AGC	CCI	GGI	CTC	CCI	CCI	'GAA	CCC	GTG	TGC	2195
713	Y	L	L	F	М	I	G	Y	A	s	A	L	V	s	L	L	N	P	С	A	732
2196																				GTG	2255
733																			S		752
2256	CCG	TGA	CAG	CGA	GAC	CTI	CAG	CAC	CTI	CCI	CCI	GG	ACC1	GT	1AT	GCI	GAC	CAT	CGG	CAT	2315
753																			G		772
2316	GGG	CGA	CCI	'GGA	GAI	GCI	GAC	CAC	CAC	CAA	\GT#	CCC	CCG	rgg	CTI	CAT	CAI	CCI	CGCI	GGT	2375
773																			L		792
2376	GAC	CT	CAI	CAI	CCI	CAC	CTC	TG	rgci	rgC1	CC:	CA	ACA?	rgc:	rca:	rTG	CCI	CA	rggg	CGA	2435
793																			G		812
2436	GAC	:AG	GGC	CCA	\GG1	CTC	CAZ	AGG!	AGAC	GCA?	AGC	ACA!	CTC	3GA	AGC:	rgcz	AGTO	GGG	CAC	CAC	2495
813				_					S											T	832
2496	CAI	CCI	:GG/	CAI	TG	GCC	CTC	CT.	rcc	CCG	TAT:	rcc:	TGA(GGA.	AGG	CCT	rcco	CT	CTGG	GGA	2555
833																			G		852
2556	GA!	rgg:	CAC	CGI	rgg	CAZ	\GA(CT	CGG	ACG(3CA	CTC	CTG	ACC	GCA	GT	GTC	CT:	CAC	GGT	2615
853																					872
2616																					
873																					
2676	CAZ	AGA.	ATG	AGAC	CTI	/CC	AGT	ATT.	ATG	GCT'	rcT(CGC	ATA	CCG	TGG(GCC	GCC:	rcc	GCA	GGA	2735
893																				D	•
2736	TC	3CT(GT(CTC	CGG:	rgg:	CAC	CCC	GCG:	TGG'	rgg	AAC'	TGA.	ACA	AGA	ACT	CGA	ACC	CGG	ACGA	2795
913																					932
2796	GG:	rgg:	rgg:	rgco	CTC:	rgg	ACA	GCA'	TGG	GGA	ACC	CCC	GCT	GCG	ATG	GCC	ACC	AGC.	AGG(STTA	2855
033	17	v	v	D	т.	D	S	M	G	N	P	Ŕ	C	D	G	н	0	0	G	Y	952

FIG. 18cont'd

2856	CCCCGCAAGTGGAGGACTGATGACGCCCCGCTCtagggactgcagcccagcccagctt	2915
953	PRKWRTDDAPL	963
2916	ctctgcccactcatttctagtccagccgcatttcagcagtgccttctggggtgtcccccc	2975
2976	acaccctgctttggccccagaggcgagggaccagtggaggtgccagggaggccccaggac	3035
3036	cctgtggtcccctggctctgcctcccaccctggggtgggggctcccggccacctgtctt	3095
3096	getectatggagteacataagecaaegecagagecectecaceteaggeeccagecectg	3155
3156	cctctccattatttatttgctctgctctcaggaagcgacgtgacccctgccccagctgga	3215
3216	acctggcagaggccttaggaccccgttccaagtgcactgcccggccaagccccagcctca	3275
3276	gcetgcgcctgagctgcatgcgccaccatttttggcagcgtggcagctttgcaaggggct	3335
3336	ggggccctcggcgtgggccatgccttctgtgtgttctgtagtgtctgggatttgccggt	3395
3396	gctcaataaatgtttattcattgaaaaaaaaaaaaaaa 3433	

FIG. 18cont'd

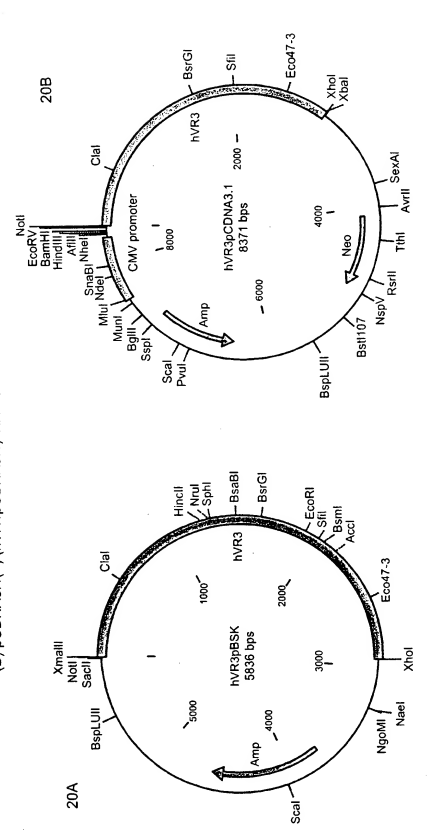
FIG. 19

AMINO ACID SEQUENCE OF hVR3

	AIVI	INO ACID S	BEQUENCE	OF HVK3	•
1	MPRVVGPGAN I	LCFQVRERGS	CCSSRLRLAA	NHIWEWPPCA	PVITTVALKQ
51	LAALLLVHVG (GGFLEPPPLA	GFCLTPLSFP	CRLSSADGPG	AGMADSSEGP
101	RAGPGEVAEL I	PGDESGTPGG	EAFPLSSLAN	LFEGEDGSLS	PSPADASRPA
151	GPGDGRPNLR 1	MKFQGAFRKG	VPNPIDLLES	TLYESSVVPG	PKKAPMDSLF
201	DYGTYRHHSS I	ONKRWRKKII	ekopospkap	APOPPPILKV	FNRPILFDIV
251	SRGSTADLDG 1	LLPFLLTHKK	RLTDEEFREP	STGKTCLPKA	LLNLSNGRND
301	TIPVLLDIAE	RTGNMREFIN	SPFRDIYYRG	QTALHIAIER	RCKHYVELLV
351	aqgadvhaqa i	RGRFFQPKDE	GGYFYFGELP	LSLAACTNOP	HIVNYLTENP
401	HKKADMRRQD	SRGNTVLHAL	VAIADNTREN	TKFVTKMYDL	LLLKCARLFP
451	DSNLEAVLNN [DGLSPLMMAA	KTGKIGIFQH	IIRREVTDED	TRHLSRKSKD
501	WAYGPVYSSL	YDLSSLDTCG	EEASVLEILV	YNSKIENRHE	MLAVEPINEL
551	LRDKWRKFGA	VSEYINVVSY.	LCAMVIETLT	AYY <mark>OPLECTP</mark>	PYPYRTTVDY
601	LRLAGEVITLE		7 - 7 - 7		
651	LVIVSAALYË	AGIEAYLAMM	VEALVLGWMN	ALYFTRGLKU	ETGTYSIMIOK
701	ILFKOLFREL	LVYLLEMIGY.	ASALVSLLNP	CANMKVCNED	QTNCTVPTYP
751	SCRDSETFST	FLLDLFKLTI	GMGDLEMLSS	TKYPVVEIIL	LVTYIILTSV
801	LLLNMLTALM	ETVGQVSKE	SKHIWKLQWA	TTILDIERSF	PVFLRKAFRS
851	GEMVTVGKSS	DGTPDRRWCF	RVDEVNWSHW	NONLGIINED	PGKNETYQYY
901	GFSHTVGRLR	RDRWSSVVPR	VVELNKNSNP	DEVVVPLDSM	GNPRCDGHQQ
951	GYPRKWRTDD	APL	•		
Ke	Y.				

 Transmer	brane	do	mains	
 Ankvrin	bindir	ng	domai	מ

FULL-LENGTH hVR3 CLONED INTO (A) pBLUESCRIPT SK(+) (hVR3pBSK) AND (B) pCDNA3.1(+) (hVR1pCDNA3.1) VIA NotI/XhoI RESTRICTION SITES. FIG. 20



SUBSTITUTE SHEET (RULE 26)

FIG. 21

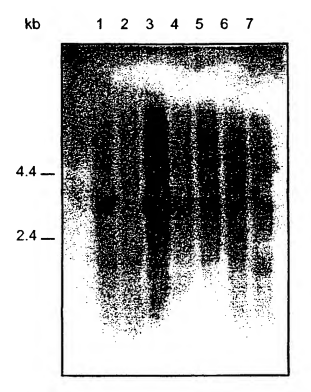
A MULTIPLE COMPARISON OF THE AMINO ACID SEQUENCES OF THE RAT VR1 AND THE HUMAN VANILLOID RECEPTORS, hVR1, hVRL-1 AND hRV3

		10	20	30	40	50
VR1	~~~~~	. ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~	~~~~~~	~~~~~~~	~~~~~~	~~~~
hVR1	~~~~~	~~~~~~~	~~~~~~	~~~~~~	~~~~~~~	~~~~
hVRL-1	~~~~~	~~~~~~	~~~~~~	~~~~~~	~~~~~~~	~~~~
hVR3	MPRVVGP	ANLCFQVRE	RGSCCSSRI	RLAANHIWE	WPPCAPVITT	VALKQ
		60	70	80	90	100
VR1	~~~~~	-~~~~~~	~~~~~~	~~~~~~	~~~~~~	~~~~
hVR1	~~~~~	.~~~~~~~	~~~~~~	~~~~~~~	~~~~~~~~	~~~~
hVRL-1	~~~~~		~~~~~~	~~~~~~	~~~~~~~	~~~~
hVR3	LAALLLVI	VGGGFLEPF	PLAGFCLTP	LSFPCRLSS.	ADGPGAGMAD:	SSEGP
		•				
. m.1		110	120	130	140 DSEESESPPO	150
VR1 hVR1					DLGAAADPLO	
hVRL-1			~~~~~~~	~~mikkwssi.	DIGAAADPIQ	KDTCP
hVR3	DACDCEV/			CIANT PECE		
IIVKS	RAGPGEVA	ELPGDE 3G1	PGGEAF PLS	SLANLFEGE.	DGSLSPSPAD.	ASKPA
		160	170	180	190	200
VR1		· · · · · · · · · · · · · · · · · · ·		Sec. 20 191	DSEEASPLDC	1 1 1 1 1 1 1 1 1
hVR1					DSEEAFPVDC	
hVRL-1					GQEDGSEADR	
hVR3	GĔĠĎĠĬĠĬ	ilrmkfogaf	RKGVPNP	IDLLES	TLYESSVVPG	PKKAP
		210	220	230	240	250
VR1	GLASCPI	TVSSVLTIC	RPGDGPASV	RPSSQDSVS	AG.EKP.PRL	YDRRS
hVR1	ELDSCET	TVSPVITÍC	RPGDGPTGA	RLLSQDSVA	ASTEKT LRL	YDRRS
hVRL-1	GSGLPPM.	. ESQFQGED	RKFAPQIRV	NLNYRKGTG	ASOPDP.NR.	FDRDR
hVR3	MDSLFDY	TYRHHSSDN	KRWRKKIIE	KQPQSPKAP	APOPPPILKV	FNRPI
		260	270	280	290	300
VR1	IFDAVAOS	NCOELESLI	PFLORSKKR		ETGKTCLLKA	MINIH
hVR1					ÉTGKTCLLKÁ	
hVRL-1					SŤGKTCLMKA	
hVR3	LFDIVSRO	STADLDGLI	PFLLTHKK	LTDEEFREP	STGKTCLPKA	LLNLS
		310	320	330	340	350
VR1	NCONDTI				QTALHIAIER	
hVR1	DGONTTI	LIJE TAROT	DSLKELVNA	SYTOSYVKO	TALHIAIER	RNMAT
hVRL-1					HSALHIAIEK	
hVR3					<u>OTALHIAIER</u>	
	SACH CANDINGS				r-accommensors	
VR1	Control Control	360	370	380	390 ESLAACTNOL	400
hVR1	VIII TOPE	ADVOAAANG	DEENVOY	PGETEGEDE	LSLAACTNOL LSLAACTNOL	ATAKE
hVRL-1					LSLAACTKOW	
hVR3					LSLAACTNOP	
114113	125 TT 14 V 5 6					
		410	420	430	440	450
VR1	LLONSWOI	ADISARDSV	GNTVLHALV	EVADNIVDN	TKFVTSMYNE	ILILG
hVR1					ikevtsmyne	
hVRL-1					IALVTSMYDG	
hVR3	LTENPHKE	ŸNWKKŐĎSK	CNTATHATA	ALADNTREN	TŘEVTKMYDL	LILKÇ
		460	470	480	490	500
VR1					ILORETHEPE	
hVR1	AKTHPTLE	LEELINKKG	MTPLALAAG	TGKŢĠVĹĂY	ILORE I QE PE	CRHLS
hVRL-1	ARICPIVO	LEDIRNLOD	LTPLKLAAK	EGKIEIFRH	ILOREFSG	LSHES,
hVR3	ARDFPDSN	ŢEAVLNNDG	ispimmäak	TCKIGIFOH	I I REEVTDED	PRITTS

	510 520 530 540 550
VR1	RKFTEWAYGPVHSSLYDLSCIDTC.EKNSVLEVIAYSSSETPNRHDMLLV
hVR1	RKFTEWAYGPVHSSLYDLSCIDTC.EKNSVLEVIAYSSSETPNRHDMLLV
hVRL-1	RKFTEWCYGPVRVSLYDLASVDSC.EENSVLEIIAF.HCKSPHRHRMVVL
hVR3	RKSKDWAYGPVYSSLYDLSSLDTCGEEASVLEILVY.WSKIENRHEMLAV
	560 570 580 590 600
VR1	EPLNRLLQDKWDRFVKRIFYFNFFVYCLYMIIFTAAAYYRPVEGLPPY
hVR1	EPLNRLLQDKWDRFVKRIFYFNFLVYCLYMIIFTMAAYYRPVDGLPPF
hVRL-1	EPLNKLLQAKWDLLIPK.FFLNFLCNLIYMFIFTAVAYHQPTLKKQAAPH
hVR1-1	EPINELLRDKWRKFGAVSFYINVVSYLCAMVIFTLTAYYQPLEGTPPY
nvks	
	610 620 630 640 650
VR1	KLKNTVGDYFRVTGE ILSVSGGVYFFFRGIQ.YFLQRRPSLKSLFVDSYS
hVR1	KMEN.IGDYFRVTGEILSVLGGVYFFFRGIQ.YFLQRRPSMKTLFVDSYS
hVRL-1	.LNAEVGNSMLLTGHILILLGGIYLLVGQLW.YFWRRHVFIWISFIDSYF
hVR3	PYRTTV.DYLRLAGEVITLFTGVLFFFTNIKDLFMKKCPGVNSLFIDGSF
	660 670 680 690 700
VR1	EILFFVQSLFMLVSVVLYFSQRKEYVASMVFSLAMGWTNMLYYTRGFQQM
hVR1	EMLFFLOSLFMLATVVLYFSHLKEYVASMVFSLALGWTNMLYYTRGFQQM
hVRL-1	EILFLFQALLTVVSQVLCFLAIEWYLPLLVSALVLGWLNLLYYTRGFQHT
hVR3	OLLYFIYSVLVIVSAALYLAGIEAYLAMMVFALVLGWMNALYFTRGLKLT
	-
VR1	GIYAVMIEKMILRDLCRFMFVYLVFLFGFSTAVVTLIEDGKNNSLP
hVR1	GIYAVMIEKMILRDLCRFMFVYIVFLFGFSTAVVTLIEDGKNDSLP
hVRL-1	GIYSVMIQKVILRDLLRFLLIYLVFLFGFAVALVSLSQEAWRPEAPTGPN
hVR3	GTYSIMIQKILFKDLFRFLLVYLLFMIGYASALVSLLNPCANMKVCNEDQ
	760 770 780 790 800
VR1	MESTPHKCRGSACK.PGNSYNSLYSTCLELFKFTIGMGDLEFTENYDFKA
hVR1	SESTSHRWRGPACRPPDSSYNSLYSTCLELFKFTIGMGDLEFTENYDFKA
hVRL-1	atesvopmegoedegngaoyrgileaslelfkftigmgelafoeolhfrg
hVR3	TNCTVPTYPSCR.DSETFSTFLLDLFKLTIGMGDLEMLSSTKYPV
	810 820 830 840 850
VR1	VFIILLLAYVILTYILLINMLIALMGETVNKIAQESKNIWKLQRAITILD
hVR1	VFIILLAYVILTYILLINMLIALMGETVNKIAQESKNIWKLQRAITILD
hVRL-1	MVLLLLAYVLLTYILLINMLIALMSETVNSVATDSWSIWKLQKAISVLE
hVR3	VFIILLVTYIILTSVLLLNMLIALMGETVGQVSKESKHIWKLQWATTILD
	860 870 880 890 900
VR1	TEKSFLKCMRKAFRSGKLLQVGFTPDGKDDYRWCFRVDEVNWTTWNTNVG
hVR1	TEKSFLKCMRKAFRSGKLLQVGYTPDGKDDYRWCFRVDEVNWTTWNTNVG
hVRL-1	MENGYWWC.RKKORAGVMLTVGTKPDGSPDERWCFRVEEVNWASWEOTLP
hVR3	IERSFPVFLRKAFRSGEMVTVGKSSDGTPDRRWCFRVDEVNWSHWNQNLG
nvks	The state of the s
	910 920 930 940 950
VR1	IINEDPGNCE GVKRTLSFSLRSG RVSGRNWKNFALV
hVR1	I INEDPGNCE GVERTLSFSLRSS RVSGRHWENFALV
hVRL-1	TLCEDPSGAGVPRTLENPVLASPPKEDEDGASEENYVPV
hVR3	IINEDPGKWETYQYYGFSHTVGRLRRDRWSSVVPRVVELNKNSNPDEVVV
	960 970 980 990
VR1	PILKDASTRORHATQQEEVQLKHYTGSLKPEDAEVFKDSMVPGEN
hVR1	PLIREASARDROSAOPEEVYLROFSGSLKPEDAEVFKSPAASGEN
hVRL-1	OLLOSN
hVR3	PLDSMGNPRCDGHQQGYPRKWRTDDAPL
J	

FIG. 21contro

FIG. 22A HYBRIDISATION OF A NORTHERN BLOT WITH hVR3



LANE 1: BONE MARROW

LANE 2: ADRENAL GLAND LANE 6: THYROID

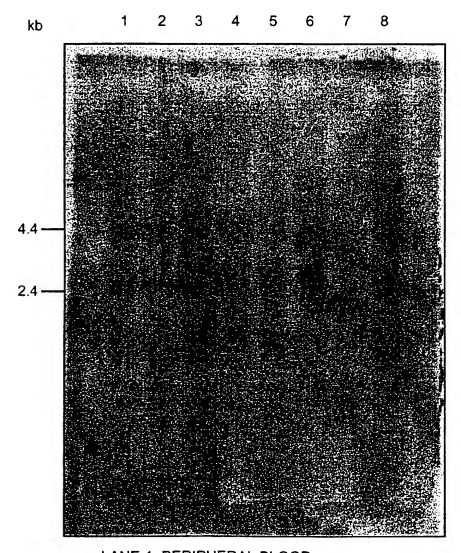
LANE 3: TRACHEA

LANE 4: LYMPH NODE

LANE 5: SPINAL CORD

LANE 7: STOMACH

FIG. 22B
HYBRIDISATION OF NORTHERN BLOT WITH hVR3 PROBE



LANE 1: PERIPHERAL BLOOD

LEUKOCYTE

LANE 2: COLON

LANE 3: SMALL INTESTINE

LANE 4: UTERUS

LANE 5: TESTIS

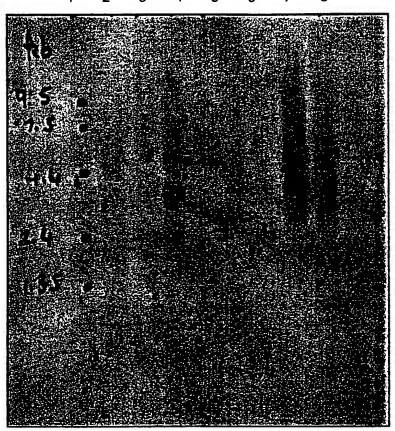
LANE 6: PROSTATE

LANE 7: THYROID

LANE 8: SPLEEN

FIG. 22C HYBRIDISATION OF A MULTI-TISSUE NORTHERN BLOT WITH THE hVR3 PROBE

1 2 6 8



LANE 1: HEART

LANE 2: BRAIN

LANE 3: PLACENTA LANE 4: LUNG

LANE 5: LIVER

LANE 6: SKELETAL MUSCLE

LANE 7: KIDNEY LANE 8: PANCREAS

SEQUENCE LISTING

<110> Glaxo Group Ltd
Tate, Simon N
Delany, Natalie S
Sanseau, P

<120> Novel Receptors

<130> PG3606

<140>

<141>

<150> GB 9826359.3

<151> 1998-12-01

<160> 40

<170> PatentIn Ver. 2.1

<210> 1

<211> 4365

<212> DNA

<213> Homo sapiens

<220>

<221> CDS

<222> (775)..(3294)

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tgtataaget cagtggetgt ggcagegagg ttgaagagea aaggeaggee gggcaeettgg 180

WO 00/32766 2 PCT/EP99/09284

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gcc	99999	gcc	tgtc	Cacc	ct c	ccag	geega	a cg	tcag	tggc	cgc	agga	ctg	cctg	ggccct	540
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ccg	gcgt	ggt (ggata	gctgo	ca g	gttg	cacac	tg:	ggcc	acag	agg	atcc	agc	aagg		777
										•					Met 1	
														ctc		825
Lys	Lys	Trp	Ser 5	Ser	Thr	Asp	Leu	Gly 10	Ala	Ala	Ala	Asp	Pro 15	Leu	Gln	
aag	gac	acc	tgc	cca	gac	ccc	ctg	gat	gga	gac	cct	aac	tcc	agg	cca	873
Lys	Asp	Thr	Cys	Pro	Asp	Pro	Leu	Asp	Gly	Asp	Pro	Asn	Ser	Arg	Pro	
		20					25					30				
cct	cca	gcc	aag	ccc	cag	ctc	tcc	acg	gcc	aag	agc	cgc	acc	cgg	ctc	921
Pro	Pro	Ala	Lys	Pro	Gln	Leu	Ser	Thr	Ala	Lys	Ser	Arg	Thr	Arg	Leu	
	35					40					45					
ttt	g gg	aag	ggt	gac	tcq	gag	gag	gct	ttc	ccq	ata	gat	tac	cct	cac	9 69

Phe Gly Lys Gly Asp Ser Glu Glu Ala Phe Pro Val Asp Cys Pro His

WO 00/32766	3	PCT/EP99/09284
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Glu	Glu	Gly	Glu	Leu	Asp	Ser	Сув	Pro	Thr	Ile	Thr	Val	Ser	Pro	Val	
				70					75					80		
atc	acc	atc	cag	agg	cca	gga	gac	ggc	ccc	acc	ggt	gcc	agg	ctg	ctg	1065
Ile	Thr	Ile	Gln	Arg	Pro	Gly	Asp	Gly	Pro	Thr	Gly	Ala	Arg	Leu	Leu	
			85					90					95			
tcc	cag	gac	tct	gtc	gcc	gcc	agc	acc	gag	aag	acc	ctc	agg	ctc	tat	1113
Ser	Gln	Asp	Ser	Val	Ala	Ala	Ser	Thr	Glu	Lys	Thr	Leu	Arg	Leu	Tyr	
		100					105					110				
	•															
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	115					120					125					
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Leu	Glu	Ser	Leu	Leu	Leu	Phe	Leu	Gln	Lув	Ser	Lys	Lys	His	Leu	Thr	
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Asp	Asn	Glu	Phe	Lys	Asp	Pro	Glu	Thr	Gly	Lys	Thr	Cys	Leu	Leu	Lys	
				150					155					160		
gcc	atg	ctc	aac	ctg	cac	gac	gga	cag	aac	acc	acc	atc	ccc	ctg	ctc	1305
Ala	Met	Leu	Asn	Leu	His	Asp	Gly	Gln	Asn	Thr	Thr	Ile	Pro	Leu	Leu	
			165					170					175			
ctg	gag	atc	gcg	cgg	caa	acg	gac	agc	ctg	aag	gag	ctt	gtc	aac	gcc	1353
Leu	Glu	Ile	Ala	Arg	Gln	Thr	Asp	Ser	Leu	Lys	Glu	Leu	Val	Asn	Ala	
		180					185					190				
agc	tac	acg	gac	agc	tac	tac	aag	ggc	cag	aca	gca	ctg	cac	atc	gcc	1401
Ser	Tyr	Thr	Asp	Ser	Tyr	Tyr	Lys	Gly	Gln	Thr	Ala	Leu	His	Ile	Ala	
	195					200					205					

															c gga	
11	e Gl	u Ar	g Ar	g As:	n Met	Ala	Leu	va]	Th	r Lei	ı Leı	ı Vai	l Gli	a Ası	n Gly	
21	0				215	,				220)				225	
gc	a gad	gt	c ca	g gc1	geg	ged	cat	999	gad	tto	: ttt	aaq	aaa	a acc	e aaa	1497
															Lys	
				230				_	235					240		
g g9	g cgg	g cct	gg:	a tto	: tac	tto	: gat	gaa	cto		cto	ı too	· ctc		gcg	1545
															Ala	1545
			245		•		2	250			Deu	Jer			. ATA	
								230					255	•		
tac	acc	: aac	: cac	cto	aac	ato	ata	220	++-						: tgg	
															Trp	1593
-,-		260		. Dec	Gly	116	265	гåя	Pne	Leu	Leu			Ser	Trp	
		200	•				205					270				
Can																
					agc											1641
GIN	275		Asp	, TI6	Ser		Arg	Asp	Ser	Val	Gly	Asn	Thr	Val	Leu	
	2/5					280					285				•	
					gtg											1689
		Leu	Val	Glu	Val	Ala	Asp	Asn	Thr	Ala	Asp	Asn	Thr	Lys	Phe	
290					295					300					305	
					aat											1737
Val	Thr	Ser	Met	Tyr	Asn	Glu	Ile	Leu	Ile	Leu	Gly	Ala	Lys	Leu	His	
				310					315					320		
ccg	acg	ctg	aag	ctg	gag	gag	ctc	acc	aac	aag	aag	gga	atg	acg	ccg	1785
					Glu											
			325					330					335			
ctg	gct	ctg	gca	gct	ggg	acc	ggg	aag	atc	ggg	gtc	tta	gcc	tat	att	1833
					Gly											
		340			-		345	-		3		350		-1-		

	cag															1881
Leu	Gln	Arg	Glu	Ile	Gln	Glu	Pro	Glu	Сув	Arg	His	Leu	Ser	Arg	Lys	
	355					360					365					
				•												
ttc	acc	gag	tgg	gcc	tac	9 99	ccc	gtg	cac	tcc	tcg	ctg	tac	gac	ctg	1929
Phe	Thr	Glu	Trp	Ala	Tyr	Gly	Pro	Val	His	Ser	Ser	Leu	Tyr	Asp	Leu	
370					375					380					385	
tcc	tgc	atc	gac	acc	tgc	gag	aag	aac	tcg	gtg	ctg	gag	gtg	atc	gcc	1977
Ser	Сув	Ile	Asp	Thr	Сув	Glu	Lys	Asn	Ser	Val	Leu	Glu	Val	Ile	Ala	
				390					395					400		
tac	agc	agc	agc	gag	acc	cct	aat	cgc	cac	gac	atg	ctc	ttg	gtg	gag	2025
Tyr	Ser	Ser	Ser	Glu	Thr	Pro	Asn	Arg	His	Asp	Met	Leu	Leu	Val	Glu	
			405					410					415			
ccg	ctg	aac	cga	ctc	ctg	cag	gac	aag	tgg	gac	aga	ttc	gtc	aag	cgc	2073
Pro	Leu	Asn	Arg	Leu	Leu	Gln	Asp	Lys	Trp	Asp	Arg	Phe	Val	Lys	Arg	
		420					425					430				
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Ile	Phe	Tyr	Phe	Asn	Phe	Leu	Val	Tyr	Сув	Leu	Tyr	Met	Ile	Ile	Phe	
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Thr	Met	Ala	Ala	Tyr	Tyr	Arg	Pro	Val	Asp	Gly	Leu	Pro	Pro	Phe	Lys	
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atg	gaa	aaa	att	gga	gac	tat	ttc	cga	gtt	act	gga	gag	atc	ctg	tct	2217
Met	Glu	Lys	Ile	Gly	Asp	Tyr	Phe	Arg	Val	Thr	Gly	Glu	Ile	Leu	Ser	
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gtg	tta	gga	gga	gtc	tac	ttc	ttt	ttc	cga	ggg	att	cag	tat	ttc	ctg	2265
Val	Leu	Gly	Gly	Val	Tyr	Phe	Phe	Phe	Arg	Gly	Ile	Gln	Tyr	Phe	Leu	
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Gln	Arg	Arg	Pro	Ser	Met	Lys	Thr	Leu	Phe	Val	Asp	Ser	Tyr	Ser	Glu	
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Met	Leu	Phe	Phe	Leu	Gln	Ser	Leu	Phe	Met	Leu	Ala	Thr	Val	Val	Leu	
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Cys	Arg	Phe	Met	Phe	Val	Tyr	Ile	Val	Phe	Leu	Phe	Gly	Phe	Ser	Thr	
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Ala	Val	Phe	Ile	Ile	Leu	Leu	Leu	Ala	Tyr	Val	Ile	Leu	Thr	Tyr	Ile	
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770	-, 5	y	4111		775	* 11E	SET	Den	wid		Sel	nrg	AGI	ser	_	
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gct cga gat agg cag tct gct cag ccc gag gaa gtt tat ctg cga cag 3225
Ala Arg Asp Arg Gln Ser Ala Gln Pro Glu Glu Val Tyr Leu Arg Gln
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ttt tca ggg tct ctg aag cca gag gac gct gag gtc ttc aag agt cct 3273
Phe Ser Gly Ser Leu Lys Pro Glu Asp Ala Glu Val Phe Lys Ser Pro
820 825 830

gcc gct tcc ggg gag aag tga ggacgtcacg cagacagcac tgtcaacact 3324
Ala Ala Ser Gly Glu Lys
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Pro Pro Pro Ala Lys Pro Gln Leu Ser Thr Ala Lys Ser Arg Thr Arg
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Leu Phe Gly Lys Gly Asp Ser Glu Glu Ala Phe Pro Val Asp Cys Pro 50 55 60

His Glu Glu Glu Leu Asp Ser Cys Pro Thr Ile Thr Val Ser Pro 65 70 75 80

Val Ile Thr Ile Gln Arg Pro Gly Asp Gly Pro Thr Gly Ala Arg Leu 85 90 95 Leu Ser Gln Asp Ser Val Ala Ala Ser Thr Glu Lys Thr Leu Arg Leu 100 105 110

Tyr Asp Arg Arg Ser Ile Phe Glu Ala Val Ala Gln Asn Asn Cys Gln 115 120 125

Asp Leu Glu Ser Leu Leu Leu Phe Leu Gln Lys Ser Lys Lys His Leu 130 135 140

Lys Ala Met Leu Asn Leu His Asp Gly Gln Asn Thr Thr Ile Pro Leu 165 170 . 175

Leu Leu Glu Ile Ala Arg Gln Thr Asp Ser Leu Lys Glu Leu Val Asn 180 185 190

Ala Ser Tyr Thr Asp Ser Tyr Tyr Lys Gly Gln Thr Ala Leu His Ile 195 200 205

Ala Ile Glu Arg Arg Asn Met Ala Leu Val Thr Leu Leu Val Glu Asn 210 215 220

Gly Ala Asp Val Gln Ala Ala Ala His Gly Asp Phe Phe Lys Lys Thr 225 230 235 240

Lys Gly Arg Pro Gly Phe Tyr Phe Gly Glu Leu Pro Leu Ser Leu Ala 245 250 255

Ala Cys Thr Asn Gln Leu Gly Ile Val Lys Phe Leu Leu Gln Asn Ser 260 265 270

Trp Gln Thr Ala Asp Ile Ser Ala Arg Asp Ser Val Gly Asn Thr Val 275 280 285

Leu His Ala Leu Val Glu Val Ala Asp Asn Thr Ala Asp Asn Thr Lys
290 295 300

Phe Val Thr Ser Met Tyr Asn Glu Ile Leu Ile Leu Gly Ala Lys Leu 305 310 315 320

His Pro Thr Leu Lys Leu Glu Glu Leu Thr Asn Lys Lys Gly Met Thr 325 330 335

Pro Leu Ala Leu Ala Ala Gly Thr Gly Lys Ile Gly Val Leu Ala Tyr 340 345 350

Ile Leu Gln Arg Glu Ile Gln Glu Pro Glu Cys Arg His Leu Ser Arg
355 360 365

Lys Phe Thr Glu Trp Ala Tyr Gly Pro Val His Ser Ser Leu Tyr Asp 370 375 380

Leu Ser Cys Ile Asp Thr Cys Glu Lys Asn Ser Val Leu Glu Val Ile 385 390 395 400

Ala Tyr Ser Ser Ser Glu Thr Pro Asn Arg His Asp Met Leu Leu Val
405 410 415

Glu Pro Leu Asn Arg Leu Leu Gln Asp Lys Trp Asp Arg Phe Val Lys
420 425 430

Arg Ile Phe Tyr Phe Asn Phe Leu Val Tyr Cys Leu Tyr Met Ile Ile 435 440 445

Phe Thr Met Ala Ala Tyr Tyr Arg Pro Val Asp Gly Leu Pro Pro Phe 450 455 460

Lys Met Glu Lys Ile Gly Asp Tyr Phe Arg Val Thr Gly Glu Ile Leu 465 470 475 480

Ser Val Leu Gly Gly Val Tyr Phe Phe Phe Arg Gly Ile Gln Tyr Phe
485 490 495

Leu Gln Arg Pro Ser Met Lys Thr Leu Phe Val Asp Ser Tyr Ser 500 505 510

Glu Met Leu Phe Ph Leu Gln Ser Leu Phe Met Leu Ala Thr Val Val 515 520 525

Leu Tyr Phe Ser His Leu Lys Glu Tyr Val Ala Ser Met Val Phe Ser 530 535 540

Leu Ala Leu Gly Trp Thr Asn Met Leu Tyr Tyr Thr Arg Gly Phe Gln 545 550 555 560

Gln Met Gly Ile Tyr Ala Val Met Ile Glu Lys Met Ile Leu Arg Asp 565 570 575

Leu Cys Arg Phe Met Phe Val Tyr Ile Val Phe Leu Phe Gly Phe Ser 580 585 590

Thr Ala Val Val Thr Leu Ile Glu Asp Gly Lys Asn Asp Ser Leu Pro
595 600 605

Ser Glu Ser Thr Ser His Arg Trp Arg Gly Pro Ala Cys Arg Pro Pro 610 615 620

Asp Ser Ser Tyr Asn Ser Leu Tyr Ser Thr Cys Leu Glu Leu Phe Lys 625 630 635 640

Phe Thr Ile Gly Met Gly Asp Leu Glu Phe Thr Glu Asn Tyr Asp Phe 645 650 655

Lys Ala Val Phe Ile Ile Leu Leu Leu Ala Tyr Val Ile Leu Thr Tyr 660 665 670

Ile Leu Leu Leu Asn Met Leu Ile Ala Leu Met Gly Glu Thr Val Asn 675 680 685

Lys Ile Ala Gln Glu Ser Lys Asn Ile Trp Lys Leu Gln Arg Ala Ile 690 695 700

Thr Ile Leu Asp Thr Glu Lys Ser Phe Leu Lys Cys Met Arg Lys Ala
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Phe Arg Ser Gly Lys Leu Leu Gln Val Gly Tyr Thr Pro Asp Gly Lys
725 730 735

Asp Asp Tyr Arg Trp Cys Phe Arg Val Asp Glu Val Asn Trp Thr Thr 740 745 750

Trp Asn Thr Asn Val Gly Ile Ile Asn Glu Asp Pro Gly Asn Cys Glu 755 760 765

Gly Val Lys Arg Thr Leu Ser Phe Ser Leu Arg Ser Ser Arg Val Ser 770 775 780

Gly Arg His Trp Lys Asn Phe Ala Leu Val Pro Leu Leu Arg Glu Ala 785 790 795 800

Ser Ala Arg Asp Arg Gln Ser Ala Gln Pro Glu Glu Val Tyr Leu Arg 805 810 815

Gln Phe Ser Gly Ser Leu Lys Pro Glu Asp Ala Glu Val Phe Lys Ser 820 825 830

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<213> Rattus norvegicus

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Pro Pro Pro Val Lys Pro His Ile Phe Thr Thr Arg Ser Arg Thr Arg
35 40 45

Leu Phe Gly Lys Gly Asp Ser Glu Glu Ala Ser Pro Leu Asp Cys Pro 50 55 60

Tyr Glu Glu Gly Gly Leu Ala Ser Cys Pro Ile Ile Thr Val Ser Ser
65 70 75 80

Val Leu Thr Ile Gln Arg Pro Gly Asp Gly Pro Ala Ser Val Arg Pro 85 90 95

Ser Ser Gln Asp Ser Val Ser Ala Gly Glu Lys Pro Pro Arg Leu Tyr
100 105 110

Asp Arg Arg Ser Ile Phe Asp Ala Val Ala Gln Ser Asn Cys Gln Glu 115 120 125

Leu Glu Ser Leu Leu Pro Phe Leu Gln Arg Ser Lys Lys Arg Leu Thr
130 135 140

Asp Ser Glu Phe Lys Asp Pro Glu Thr Gly Lys Thr Cys Leu Leu Lys
145 150 155 160

Ala Met Leu Asn Leu His Asn Gly Gln Asn Asp Thr Ile Ala Leu Leu 165 170 175

Leu Asp Val Ala Arg Lys Thr Asp Ser Leu Lys Gln Phe Val Asn Ala 180 185 190

Ser Tyr Thr Asp Ser Tyr Tyr Lys Gly Gln Thr Ala Leu His Ile Ala 195 200 . 205

Ile Glu Arg Arg Asn Met Thr Leu Val Thr Leu Val Glu Asn Gly
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Ala Asp Val Gln Ala Ala Ala Asn Gly Asp Phe Phe Lys Lys Thr Lys 225 230 235 240

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- Cys Thr Asn Gln Leu Ala Ile Val Lys Phe Leu Leu Gln Asn Ser Trp
 260 265 270
- Gln Pro Ala Asp Ile Ser Ala Arg Asp Ser Val Gly Asn Thr Val Leu 275 280 285
- His Ala Leu Val Glu Val Ala Asp Asn Thr Val Asp Asn Thr Lys Phe 290 295 300
- Val Thr Ser Met Tyr Asn Glu Ile Leu Ile Leu Gly Ala Lys Leu His 305 310 315 320
- Pro Thr Leu Lys Leu Glu Glu Ile Thr Asn Arg Lys Gly Leu Thr Pro 325 330 335
- Leu Ala Leu Ala Ala Ser Ser Gly Lys Ile Gly Val Leu Ala Tyr Ile 340 345 350
- Leu Gln Arg Glu Ile His Glu Pro Glu Cys Arg His Leu Ser Arg Lys 355 360 365
- Phe Thr Glu Trp Ala Tyr Gly Pro Val His Ser Ser Leu Tyr Asp Leu 370 375 380
- Ser Cys Ile Asp Thr Cys Glu Lys Asn Ser Val Leu Glu Val Ile Ala 385 390 395 400
- Tyr Ser Ser Ser Glu Thr Pro Asn Arg His Asp Met Leu Leu Val Glu
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 410
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- Pro Leu Asn Arg Leu Leu Gln Asp Lys Trp Asp Arg Phe Val Lys Arg
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- Ile Phe Tyr Phe Asn Phe Phe Val Tyr Cys Leu Tyr Met Ile Ile Phe
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Thr Ala Ala Ala Tyr Tyr Arg Pro Val Glu Gly Leu Pro Pro Tyr Lys
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Leu Lys Asn Thr Val Gly Asp Tyr Phe Arg Val Thr Gly Glu Ile Leu 465 470 475 480

Ser Val Ser Gly Gly Val Tyr Phe Phe Phe Arg Gly Ile Gln Tyr Phe
485 490 495

Leu Gln Arg Arg Pro Ser Leu Lys Ser Leu Phe Val Asp Ser Tyr Ser 500 505 510

Glu Ile Leu Phe Phe Val Gln Ser Leu Phe Met Leu Val Ser Val Val 515 520 525

Leu Tyr Phe Ser Gln Arg Lys Glu Tyr Val Ala Ser Met Val Phe Ser 530 540

Leu Ala Met Gly Trp Thr Asn Met Leu Tyr Tyr Thr Arg Gly Phe Gln 545 550 555 560

Gln Met Gly Ile Tyr Ala Val Met Ile Glu Lys Met Ile Leu Arg Asp 565 570 575

Leu Cys Arg Phe Met Phe Val Tyr Leu Val Phe Leu Phe Gly Phe Ser 580 585 590

Thr Ala Val Val Thr Leu Ile Glu Asp Gly Lys Asn Asn Ser Leu Pro 595 600 605

Met Glu Ser Thr Pro His Lys Cys Arg Gly Ser Ala Cys Lys Pro Gly 610 615 620

Asn Ser Tyr Asn Ser Leu Tyr Ser Thr Cys Leu Glu Leu Phe Lys Phe 625 630 635 640

Thr Ile Gly Met Gly Asp Leu Glu Phe Thr Glu Asn Tyr Asp Phe Lys
645 650 655

Ala Val Phe Ile Ile Leu Leu Leu Ala Tyr Val Ile Leu Thr Tyr Ile
660 665 670

Leu Leu Leu Asn Met Leu Ile Ala Leu Met Gly Glu Thr Val Asn Lys
675 680 685

Ile Ala Gln Glu Ser Lys Asn Ile Trp Lys Leu Gln Arg Ala Ile Thr
690 695 700

Ile Leu Asp Thr Glu Lys Ser Phe Leu Lys Cys Met Arg Lys Ala Phe 705 710 715 720

Arg Ser Gly Lys Leu Gln Val Gly Phe Thr Pro Asp Gly Lys Asp
725 730 735

Asp Tyr Arg Trp Cys Phe Arg Val Asp Glu Val Asn Trp Thr Thr Trp 740 745 750

Asn Thr Asn Val Gly Ile Ile Asn Glu Asp Pro Gly Asn Cys Glu Gly
755 760 765

Val Lys Arg Thr Leu Ser Phe Ser Leu Arg Ser Gly Arg Val Ser Gly 770 780

Arg Asn Trp Lys Asn Phe Ala Leu Val Pro Leu Leu Arg Asp Ala Ser 785 790 795 800

Thr Arg Asp Arg His Ala Thr Gln Gln Glu Glu Val Gln Leu Lys His 805 810 815

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Net Pro Arg Val Val Gly Pro Gly Ala

5

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Leu	Arg	Leu	Ala	Ala	Asn	His	Ile	Trp	Glu	Trp	Pro	Pro	Сув	Ala	Pro	
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gtc	att	aca	acg	gtg	gct	ttg	aag	cag	ctg	gca	gca	ctg	ctg	ctt	gtc	856
Val	Ile	Thr	Thr	Val	Ala	Leu	Lys	Gln	Leu	Ala	Ala	Leu	Leu	Leu	Val	
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cta	200															
		cac														1528
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		Lys														1576
	,		285	~1 0	Jeu		-75	290	Pan	₽Gſ	ABN		Ser 295	ASN	GIÅ	
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ege aac gac ace ate cet gtg etg etg gac ate geg gag ege ace gge Arg Asn Asp Thr Ile Pro Val Leu Leu Asp Ile Ala Glu Arg Thr Gly aac atg cgg gag ttc att aac tcg ccc ttc cgt gac atc tac tat cga Asn Met Arg Glu Phe Ile Asn Ser Pro Phe Arg Asp Ile Tyr Tyr Arg ggt cag aca gcc ctg cac atc gcc att gag cgt cgc tgc aaa cac tac Gly Gln Thr Ala Leu His Ile Ala Ile Glu Arg Arg Cys Lys His Tyr gtg gaa ctt ctc gtg gcc cag gga gct gat gtc cac gcc cag gcc cgt Val Glu Leu Leu Val Ala Gln Gly Ala Asp Val His Ala Gln Ala Arg ggg cgc ttc ttc cag ccc aag gat gag ggg ggc tac ttc tac ttt ggg Gly Arg Phe Phe Gln Pro Lys Asp Glu Gly Gly Tyr Phe Tyr Phe Gly gag ctg ccc ctg tcg ctg gct gcc tgc acc aac cag ccc cac att gtc Glu Leu Pro Leu Ser Leu Ala Ala Cys Thr Asn Gln Pro His Ile Val aac tac ctg acg gag aac ccc cac aag aag gcg gac atg cgg cgc cag Asn Tyr Leu Thr Glu Asn Pro His Lys Lys Ala Asp Met Arg Arg Gln gac tog cga ggc aac aca gtg ctg cat gcg ctg gtg gcc att gct gac Asp Ser Arg Gly Asn Thr Val Leu His Ala Leu Val Ala Ile Ala Asp aac acc cgt gag aac acc aag ttt gtt acc aag atg tac gac ctg ctg Asn Thr Arg Glu Asn Thr Lys Phe Val Thr Lys Met Tyr Asp Leu Leu

ctg	ctc	aag	tgt	gcc	cgc	ctc	ttc	ccc	gac	agc	aac	ctg	gag	gcc	gtg	2056
Leu	Leu	Lys	Сув	Ala	Arg	Leu	Phe	Pro	Asp	Ser	Asn	Leu	Glu	Ala	Val	
			445					450					455			
ctc	aac	aac	gac	ggc	ctc	tcg	ccc	ctc	atg	atg	gct	gcc	aag	acg	ggc	2104
Leu	Asn	Asn	Авр	Gly	Leu	Ser	Pro	Leu	Met	Met	Ala	Ala	Lув	Thr	Gly	
		460					465					470				
aag	att	g gg	atc	ttt	cag	cac	atc	atc	cgg	cgg	gag	gtg	acg	gat	gag	2152
Lys	Ile	Gly	Ile	Phe	Gln	His	Ile	Ile	Arg	Arg	Glu	Val	Thr	Asp	Glu	
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gac	aca	cgg	cac	ctg	tcc	cgc	aag	tcc	aag	gac	tgg	gcc	tat	ggg	cca	2200
Asp	Thr	Arg	His	Leu	Ser	Arg	Lys	Ser	Lys	Asp	Trp	Ala	Tyr	Gly	Pro	
490					495					500					505	
gtg	tat	tcc	tcg	ctt	tat	gac	ctc	tcc	tcc	ctg	gac	acg	tgt	ggg	gaa	2248
Val	Tyr	Ser	Ser	Leu	Tyr	Asp	Leu	Ser	Ser	Leu	Asp	Thr	Cys	Gly	Glu	
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gag	gcc	tcc	gtg	ctg	gag	atc	ctg	gtg	tac	aac	agc	aag	att	gag	aac	2296
Glu	Ala	Ser	Val	Leu	Glu	Ile	Leu	Val	Tyr	Asn	Ser	Lys	Ile	Glu	Asn	
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cgc	cac	gag	atg	ctg	gct	gtg	gag	CCC	atc	aat	gaa	ctg	ctg	cgg	gac	2344
Arg	His	Glu	Met	Leu	Ala	Val	Glu	Pro	Ile	Asn	Glu	Leu	Leu	Arg	Asp	
		540					545					550				
aag	tgg	cgg	aag	ttc	9 99	gcc	gtc	tcc	ttc	tac	atc	aac	gtg	gtc	tcc	2392
Lys	Trp	Arg	Lys	Phe	Gly	Ala	Val	Ser	Phe	Tyr	Ile	Asn	Val	Val	Ser	
•	555					560					565					
			gcc													2440
Tyr	Leu	Сув	Ala	Met	Val	Ilè	Phe	Thr	Leu	Thr	Ala	Tyr	Tyr	Gln	Pro	
570					5 75					580					5 85	

ctq qaq ggc aca ccg ccg tac cct tac cgc acc acg gtg gac tac ctq Leu Glu Gly Thr Pro Pro Tyr Pro Tyr Arg Thr Thr Val Asp Tyr Leu egg etg get gge gag gte att acg etc tte act ggg gte etg tte tte Arg Leu Ala Gly Glu Val Ile Thr Leu Phe Thr Gly Val Leu Phe Phe ttc acc aac atc aaa gac ttg ttc atg aag aaa tgc cct gga gtg aat Phe Thr Asn Ile Lys Asp Leu Phe Met Lys Lys Cys Pro Gly Val Asn tet etc ttc att gat ggc tcc ttc cag etg etc tac ttc atc tac tct Ser Leu Phe Ile Asp Gly Ser Phe Gln Leu Leu Tyr Phe Ile Tyr Ser gtc ctg gtg atc gtc tca gca gcc ctc tac ctg gca ggg atc gag gcc Val Leu Val Ile Val Ser Ala Ala Leu Tyr Leu Ala Gly Ile Glu Ala tac ctg gcc atg atg gtc ttt gcc ctg gtc ctg ggc tgg atg aat gcc Tyr Leu Ala Met Met Val Phe Ala Leu Val Leu Gly Trp Met Asn Ala ctt tac ttc acc cgt ggg ctg aag ctg acg ggg acc tat agc atc atg Leu Tyr Phe Thr Arg Gly Leu Lys Leu Thr Gly Thr Tyr Ser Ile Met atc cag aag att ctc ttc aag gac ctt ttc cga ttc ctg ctc gtc tac Ile Gln Lys Ile Leu Phe Lys Asp Leu Phe Arg Phe Leu Leu Val Tyr ttg ctc ttc atg atc ggc tac gct tca gcc ctg gtc tcc ctc ctg aac Leu Leu Phe Met Ile Gly Tyr Ala Ser Ala Leu Val Ser Leu Leu Asn

cca	tgt	gee	aac	atg	aag	ata	tgc	aat	gag	gac	cag	acc	aac	tac	aca	2920
					Lys											
730					735		•			740				•	745	
gtg	ccc	act	tac	ccc	tcg	tgc	cgt	gac	agc	gag	acc	ttc	agc	acc	ttc	2968
Val	Pro	Thr	Tyr	Pro	Ser	Сув	Arg	Asp	Ser	Glu	Thr	Phe	Ser	Thr	Phe	
				750					755					760		
ctc	ctg	gac	ctg	ttt	aag	ctg	acc	atc	ggc	atg	gġc	gac	ctg	gag	atg	3016
Leu	Leu	Asp	Leu	Phe	Lys	Leu	Thr	Ile	Gly	Met	Gly	Asp	Leu	Glu	Met	
i			765					770					775			
ctg	agc	agc	acc	aag	tac	ccc	gtg	gtc	ttc	atc	atc	ctg	ctg	gtg	acc	3064
Leu	Ser	Ser	Thr	Lys	Tyr	Pro	Val	Val	Phe	Ile	Ile	Leu	Leu	Val	Thr	
		780					785					790				
						•										
tac	atc	atc	ctc	acc	tct	gtg	ctg	ctc	ctc	aac	atg	ctc	att	gcc	ctc	3112
Tyr	Ile	Ile	Leu	Thr	Ser	Val	Leu	Leu	Leu	Asn	Met	Leu	Ile	Ala	Leu	
	795					800					805					
					ggc						_					3160
	Gly	Glu	Thr	Val	Gly	Gln	Val	Ser	Lys	Glu	Ser	Lув	His	Ile	Trp	
810					815					820					825	
					acc						-	_				3208
Lys	Leu	Gln	Trp		Thr	Thr	Ile	Leu		Ile	Glu	Arg	Ser	Phe	Pro	
				830					835					840		
					gcc											3256
vai	Pne	Leu		Lys	Ala	Phe	Arg		Gly	Glu	Met	Val		Val	Gly	
			845					850					855			
		.			+								•			
					act											3304
гла	ser		Asp	GIA	Thr	Pro		Arg	Arg	Trp	Сув		Arg	Val	Asp	
		860					865					870				

gag gtg aac	tgg tct cac	tgg aac cag	aac ttg ggc atc	atc aac gag 3352
Glu Val Asn	Trp Ser His	Trp Asn Gln	Asn Leu Gly Ile	Ile Asn Glu
875		880	885	

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Asp	Pro	Gly	Lys	Asn	Glu	Thr	Tyr	Gln	Tyr	Tyr	Gly	Phe	Ser	His	Thr	
890					895					900					905	

gtg	ggc	cgc	ctc	cgc	agg	gat	cgc	tgg	tcc	tcg	gtg	gta	ccc	cgc	gtg	3448
Val	Gly	Arg	Leu	Arg	Arg	Asp	Arg	Trp	Ser	Ser	Val	Val	Pro	Arg	Val	
				910					915					920		

gtg	gaa	ctg	aac	aag	aac	tcg	aac	ccg	gac	gag	gtg	gtg	gtg	cct	ctg	3496
Val	Glu	Leu	Asn	Lys	Asn	Ser	Asn	Pro	Asp	Glu	Val	Val	Val	Pro	Leu	
			925					930					935			

gac	agc	atg	g gg	aac	ccc	cgc	tgc	gat	ggc	cac	cag	cag	ggt	tac	CCC	3544
Asp	Ser	Met	Gly	Asn	Pro	Arg	Сув	Asp	Gly	His	Gln	Gln	Gly	Tyr	Pro	
		940					945					950				

cgc aag tgg agg act gat gac gcc ccg ctc tag ggactgcagc ccagccccag 3597
Arg Lys Trp Arg Thr Asp Asp Ala Pro Leu
955 960

cetactetgee caeteatte tagtecagee geattecage agtgeettet ggggtgteee 3657
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4118

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<211> 963

<212> PRT

<213> Homo sapiens

<400> 5

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Glu Arg Gly Ser Cys Cys Ser Ser Arg Leu Arg Leu Ala Ala Asn His
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Ile Trp Glu Trp Pro Pro Cys Ala Pro Val Ile Thr Thr Val Ala Leu
35 40 45

Lys Gln Leu Ala Ala Leu Leu Leu Val His Val Gly Gly Phe Leu 50 55 60

Glu Pro Pro Pro Leu Ala Gly Phe Cys Leu Thr Pro Leu Ser Phe Pro 65 70 75 80

Cys Arg Leu Ser Ser Ala Asp Gly Pro Gly Ala Gly Met Ala Asp Ser 85 90 95

Ser Glu Gly Pro Arg Ala Gly Pro Gly Glu Val Ala Glu Leu Pro Gly 100 105 110

Asp Glu Ser Gly Thr Pro Gly Gly Glu Ala Phe Pro Leu Ser Ser Leu 115 120 125

Ala Asn Leu Phe Glu Gly Glu Asp Gly Ser Leu Ser Pro Ser Pro Ala 130 135 140

WO 00/32766 Asp Ala Ser Arg Pro Ala Gly Pro Gly Asp Gly Arg Pro Asn Leu Arg Met Lys Phe Gln Gly Ala Phe Arg Lys Gly Val Pro Asn Pro Ile Asp Leu Leu Glu Ser Thr Leu Tyr Glu Ser Ser Val Val Pro Gly Pro Lys Lys Ala Pro Met Asp Ser Leu Phe Asp Tyr Gly Thr Tyr Arg His His Ser Ser Asp Asn Lys Arg Trp Arg Lys Lys Ile Ile Glu Lys Gln Pro Gln Ser Pro Lys Ala Pro Ala Pro Gln Pro Pro Pro Ile Leu Lys Val Phe Asn Arg Pro Ile Leu Phe Asp Ile Val Ser Arg Gly Ser Thr Ala Asp Leu Asp Gly Leu Leu Pro Phe Leu Leu Thr His Lys Lys Arg Leu Thr Asp Glu Glu Phe Arg Glu Pro Ser Thr Gly Lys Thr Cys Leu Pro Lys Ala Leu Leu Asn Leu Ser Asn Gly Arg Asn Asp Thr Ile Pro Val Leu Leu Asp Ile Ala Glu Arg Thr Gly Asn Met Arg Glu Phe Ile Asn Ser Pro Phe Arg Asp Ile Tyr Tyr Arg Gly Gln Thr Ala Leu His Ile

Ala Ile Glu Arg Arg Cys Lys His Tyr Val Glu Leu Leu Val Ala Gln

Gly Ala Asp Val His Ala Gln Ala Arg Gly Arg Phe Phe Gln Pro Lys Asp Glu Gly Gly Tyr Phe Tyr Phe Gly Glu Leu Pro Leu Ser Leu Ala Ala Cys Thr Asn Gln Pro His Ile Val Asn Tyr Leu Thr Glu Asn Pro His Lys Lys Ala Asp Met Arg Arg Gln Asp Ser Arg Gly Asn Thr Val Leu His Ala Leu Val Ala Ile Ala Asp Asn Thr Arg Glu Asn Thr Lys Phe Val Thr Lys Met Tyr Asp Leu Leu Leu Leu Lys Cys Ala Arg Leu Phe Pro Asp Ser Asn Leu Glu Ala Val Leu Asn Asn Asp Gly Leu Ser Pro Leu Met Met Ala Ala Lys Thr Gly Lys Ile Gly Ile Phe Gln His Ile Ile Arg Arg Glu Val Thr Asp Glu Asp Thr Arg His Leu Ser Arg Lys Ser Lys Asp Trp Ala Tyr Gly Pro Val Tyr Ser Ser Leu Tyr Asp Leu Ser Ser Leu Asp Thr Cys Gly Glu Glu Ala Ser Val Leu Glu Ile Leu Val Tyr Asn Ser Lys Ile Glu Asn Arg His Glu Met Leu Ala Val

Glu Pro Ile Asn Glu Leu Leu Arg Asp Lys Trp Arg Lys Phe Gly Ala

Val Ser Phe Tyr Ile Asn Val Val Ser Tyr Leu Cys Ala Met Val Ile 565 570 575

Phe Thr Leu Thr Ala Tyr Tyr Gln Pro L u Glu Gly Thr Pro Pro Tyr 580 585 590

Pro Tyr Arg Thr Thr Val Asp Tyr Leu Arg Leu Ala Gly Glu Val Ile
595 600 605

Thr Leu Phe Thr Gly Val Leu Phe Phe Phe Thr Asn Ile Lys Asp Leu 610 615 620

Phe Met Lys Lys Cys Pro Gly Val Asn Ser Leu Phe Ile Asp Gly Ser 625 630 635 640

Phe Gln Leu Leu Tyr Phe Ile Tyr Ser Val Leu Val Ile Val Ser Ala 645 650 655

Ala Leu Tyr Leu Ala Gly Ile Glu Ala Tyr Leu Ala Met Met Val Phe 660 665 670

Ala Leu Val Leu Gly Trp Met Asn Ala Leu Tyr Phe Thr Arg Gly Leu 675 680 685

Lys Leu Thr Gly Thr Tyr Ser Ile Met Ile Gln Lys Ile Leu Phe Lys 690 695 700

Asp Leu Phe Arg Phe Leu Leu Val Tyr Leu Leu Phe Met Ile Gly Tyr 705 710 715 720

Ala Ser Ala Leu Val Ser Leu Leu Asn Pro Cys Ala Asn Met Lys Val 725 730 735

Cys Asn Glu Asp Gln Thr Asn Cys Thr Val Pro Thr Tyr Pro Ser Cys
740 745 750

Arg Asp Ser Glu Thr Phe Ser Thr Phe Leu Leu Asp Leu Phe Lys Leu
755 760 765

Thr Ile Gly Met Gly Asp Leu Glu Met Leu Ser Ser Thr Lys Tyr Pro
770 775 780

Val Val Phe Ile Ile Leu Leu Val Thr Tyr Ile Ile Leu Thr Ser Val 785 790 795 800

Leu Leu Leu Asn Met Leu Ile Ala Leu Met Gly Glu Thr Val Gly Gln 805 810 815

Val Ser Lys Glu Ser Lys His Ile Trp Lys Leu Gln Trp Ala Thr Thr 820 825 830

Ile Leu Asp Ile Glu Arg Ser Phe Pro Val Phe Leu Arg Lys Ala Phe 835 840 845

Arg Ser Gly Glu Met Val Thr Val Gly Lys Ser Ser Asp Gly Thr Pro 850 855 860

Asp Arg Arg Trp Cys Phe Arg Val Asp Glu Val Asn Trp Ser His Trp 865 870 875 880

Asn Gln Asn Leu Gly Ile Ile Asn Glu Asp Pro Gly Lys Asn Glu Thr 885 890 895

Tyr Gln Tyr Tyr Gly Phe Ser His Thr Val Gly Arg Leu Arg Arg Asp 900 905 910

Arg Trp Ser Ser Val Val Pro Arg Val Val Glu Leu Asn Lys Asn Ser 915 920 925

Asn Pro Asp Glu Val Val Pro Leu Asp Ser Met Gly Asn Pro Arg 930 935 940

Cys Asp Gly His Gln Gln Gly Tyr Pro Arg Lys Trp Arg Thr Asp Asp 945 950 955 960

Ala Pro Leu

<211> 764

<212> PRT

<213> Homo sapiens

<400> 6

Met Thr Ser Pro Ser Ser Ser Pro Val Phe Arg Leu Glu Thr Leu Asp

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Gly Gly Gln Glu Asp Gly Ser Glu Ala Asp Arg Gly Lys Leu Asp Phe
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Gly Ser Gly Leu Pro Pro Met Glu Ser Gln Phe Gln Gly Glu Asp Arg
35 40 45

Lys Phe Ala Pro Gln Ile Arg Val Asn Leu Asn Tyr Arg Lys Gly Thr
50 55 60

Gly Ala Ser Gln Pro Asp Pro Asn Arg Phe Asp Arg Asp Arg Leu Phe
65 70 75 80

Asn Ala Val Ser Arg Gly Val Pro Glu Asp Leu Ala Gly Leu Pro Glu 85 90 95

Tyr Leu Ser Lys Thr Ser Lys Tyr Leu Thr Asp Ser Glu Tyr Thr Glu
100 105 \ 110

Gly Ser Thr Gly Lys Thr Cys Leu Met Lys Ala Val Leu Asn Leu Lys 115 120 125

Asp Gly Val Asn Ala Cys Ile Leu Pro Leu Leu Gln Ile Asp Arg Asp 130 135 140

Ser Gly Asn Pro Gln Pro Leu Val Asn Ala Gln Cys Thr Asp Asp Tyr 145 150 155 160

Tyr Arg Gly His Ser Ala Leu His Ile Ala Ile Glu Lys Arg Ser Leu
165 170 175

Gln Cys Val Lys Leu Leu Val Glu Asn Gly Ala Asn Val His Ala Arg 180 185 190

- Ala Cys Gly Arg Phe Phe Gln Lys Gly Gln Gly Thr Cys Phe Tyr Phe 195 200 205
- Gly Glu Leu Pro Leu Ser Leu Ala Ala Cys Thr Lys Gln Trp Asp Val 210 215 220
- Val Ser Tyr Leu Leu Glu Asn Pro His Gln Pro Ala Ser Leu Gln Ala 225 230 235 240
- Thr Asp Ser Gln Gly Asn Thr Val Leu His Ala Leu Val Met Ile Ser 245 250 255
- Asp Asn Ser Ala Glu Asn Ile Ala Leu Val Thr Ser Met Tyr Asp Gly 260 265 270
- Leu Leu Gln Ala Gly Ala Arg Leu Cys Pro Thr Val Gln Leu Glu Asp 275 280 285
- Ile Arg Asn Leu Gln Asp Leu Thr Pro Leu Lys Leu Ala Ala Lys Glu 290 295 300
- Gly Lys Ile Glu Ile Phe Arg His Ile Leu Gln Arg Glu Phe Ser Gly
 305 310 315 320
- Leu Ser His Leu Ser Arg Lys Phe Thr Glu Trp Cys Tyr Gly Pro Val
- Arg Val Ser Leu Tyr Asp Leu Ala Ser Val Asp Ser Cys Glu Glu Asn 340 345 350
- Ser Val Leu Glu Ile Ile Ala Phe His Cys Lys Ser Pro His Arg His 355 360 365
- Arg Met Val Val Leu Glu Pro Leu Asn Lys Leu Leu Gln Ala Lys Trp 370 375 380

Asp Leu Leu Ile Pro Lys Phe Phe Leu Asn Phe Leu Cys Asn Leu Ile 385 390 395 400

Tyr Met Phe Ile Phe Thr Ala Val Ala Tyr His Gln Pro Thr Leu Lys
405 410 415

Lys Gln Ala Ala Pro His Leu Lys Ala Glu Val Gly Asn Ser Met Leu
420 425 430

Leu Thr Gly His Ile Leu Ile Leu Leu Gly Gly Ile Tyr Leu Leu Val
435
440
445

Gly Gln Leu Trp Tyr Phe Trp Arg Arg His Val Phe Ile Trp Ile Ser 450 455 460

Phe Ile Asp Ser Tyr Phe Glu Ile Leu Phe Leu Phe Gln Ala Leu Leu 465 470 475 480

Thr Val Val Ser Gln Val Leu Cys Phe Leu Ala Ile Glu Trp Tyr Leu
485 490 495

Pro Leu Leu Val Ser Ala Leu Val Leu Gly Trp Leu Asn Leu Leu Tyr
500 505 510

Tyr Thr Arg Gly Phe Gln His Thr Gly Ile Tyr Ser Val Met Ile Gln
515 520 525

Lys Val Ile Leu Arg Asp Leu Leu Arg Phe Leu Leu Ile Tyr Leu Val 530 535 540

Phe Leu Phe Gly Phe Ala Val Ala Leu Val Ser Leu Ser Gln Glu Ala 545 550 555 560

Trp Arg Pro Glu Ala Pro Thr Gly Pro Asn Ala Thr Glu Ser Val Gln
565 570 575

Pro Met Glu Gly Gln Glu Asp Glu Gly Asn Gly Ala Gln Tyr Arg Gly 580 585 590

 11e
 Leu
 Glu
 Ala
 Ser
 Leu
 Glu
 Leu
 Phe
 Lys
 Phe
 Thr
 Ile
 Gly
 Met
 Gly

 Glu
 Leu
 Ala
 Phe
 Gln
 Gln
 Leu
 His
 Phe
 Arg
 Gly
 Met
 Val
 Leu
 Leu

 610
 Glo
 Fix
 Fix
 Thr
 Tyr
 Fix
 F

Leu Ile Ala Leu Met Ser Glu Thr Val Asn Ser Val Ala Thr Asp Ser 645 650 655

Trp Ser Ile Trp Lys Leu Gln Lys Ala Ile Ser Val Leu Glu Met Glu 660 665 670

Asn Gly Tyr Trp Trp Cys Arg Lys Lys Gln Arg Ala Gly Val Met Leu 675 680 685

Thr Val Gly Thr Lys Pro Asp Gly Ser Pro Asp Glu Arg Trp Cys Phe
690 695 700

Arg Val Glu Glu Val Asn Trp Ala Ser Trp Glu Gln Thr Leu Pro Thr
705 710 715 720

Leu Cys Glu Asp Pro Ser Gly Ala Gly Val Pro Arg Thr Leu Glu Asn
725 730 735

Pro Val Leu Ala Ser Pro Pro Lys Glu Asp Glu Asp Gly Ala Ser Glu
740 745 750

Glu Asn Tyr Val Pro Val Gln Leu Leu Gln Ser Asn 755 760 WO 00/32766 35 PCT/EP99/09284

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18

<210> 8

<211> 20

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: Primer

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<211> 19

<212> DNA

<213> Artificial Sequence

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<223> Description of Artificial Sequence: Primer

<400> 9

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WO 00/32766 36 PCT/EP99/09284 <210> 10 <211> 17 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence: Primer <400> 10 gtaaaacgac ggccagt 17 <210> 11 <211> 20 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence: Primer <400> 11 aattaaccct cactaaaggg 20 <210> 12 <211> 20 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence: Primer <400> 12

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37
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 WO 00/32766
<210> 13
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                                                                    20
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<400> 14
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ctgcagaact cctggcaga
<210> 15
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20

<223> Description of Artificial Sequence: Primer

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WO 00/32766 38 PCT/EP99/09284 <210> 16 <211> 21 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence: Primer <400> 16 tcctctggct tccaacccgt t 21 <210> 17 <211> 20 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence: Primer

<400> 17
gaactgggca gaaagtgcct 20

<210> 18
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ctggagttag ggtctccatc c

<211> 43

<212> DNA

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<223> Description of Artificial Sequence: Primer

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43

<210> 20

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<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: Primer

<400> 20

aggeceacte ggtgaactte

20

<210> 21

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<223> Description of Artificial Sequence: Primer

<400> 21

gacgagcatg tacaatgaga

<211> 20

<212> DNA

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<220>

<223> Description of Artificial Sequence: Primer

<400> 22

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<210> 23

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<212> DNA

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<400> 23

tgtggacagc tacagtgaga

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<210> 24

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<212> DNA

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<400> 24

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<211> 20

<212> DNA

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<220>

<223> Description of Artificial Sequence: Primer

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<210> 26

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<212> DNA

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<400> 26

gtggaaaacc cgaacaaga

19

<210> 27

<211> 23

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: Synthetic sequence

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Cys His Ile Phe Thr Thr Arg Ser Arg Thr Arg Leu Phe Gly Lys Gly

1 5 10 15

Asp Ser Glu Glu Ala Ser Cys

<211> 21

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: Synthetic sequence

<400> 28

Cys Gly Ser Leu Lys Pro Glu Asp Ala Glu Val Phe Lys Asp Ser Met

1 5 10 15

20

Val Pro Gly Glu Lys

<210> 29

<211> 20

<212> DNA

<213> Artificial Sequence

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<223> Description of Artificial Sequence: Primer

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<210> 30

<211> 18

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: Primer

<400> 30

totgocaggt tocagetg 18

<210> 31

<211> 41.

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: Primer

<400> 31

gtcatagogg cogogocoa coatgocoag ggtagttgga c

<210> 32

<211> 20

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: Primer

<400> 32

cacctcttgt tgtcactgga

<210> 33

<211> 23

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: Primer

<400> 33

caaatctgcg catgaagttc cag

23

41

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WO 00/32766
                                      45
                                                             PCT/EP99/09284
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atctcgtggc ggttctcaat

int. .tional Application No

CO7K16/28

C12N5/10

10 to

PCT/EP 99/09284 A. CLASSIFICATION OF SUBJECT MATTER
IPC 7 C12N15/12 C07K14/705 C12N15/85

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols) IPC 7 C12N C07K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

C. DOCUM	ENTS CONSIDERED TO BE RELEVANT	
Category 3	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	CATERINA, M.J. ET AL.: "The capsaicin receptor: a heat-actvated ion channel in the pain pathway" NATURE, vol. 389, no. 6653, 23 October 1997 (1997-10-23), pages 816-824, XP002075020 cited in the application abstract page 819; figures 5A,C page 820, column 2, line 13 -page 821, column 1, line 29	1-3,6,9, 14-16, 45-47
X A	page 823, column 2, line 13 - line 14 page 817, column 2, line 12 -page 820, column 1, line 21 page 823, column 2, line 19 -page 824, column 1, line 5	26 4,5,7,8, 10-13, 17-25, 48-51

Further documents are listed in the continuation of box C.	Patent family members are listed in annex.
* Special categories of cited documents :	
"A" document defining the general state of the lart which is not considered to be of particular relevance	'T' later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention.
"E" earlier document but published on or after the international filling date	"X" document of particular relevance; the claimed invention
"L" document which may throw doubts on priority claim(s) or	cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
which is cited to establish the publication date at another citation or other special reason (as specified)	"Y" document of particular relevance; the claimed invention
"O" document referring to an oral disclosure, use, exhibition or	cannot be considered to involve an inventive step when the document is combined with one or more other such docu-
other means "P" document published prior to the <i>intern</i> ational filing date but	ments, such combination being obvious to a person skilled in the art.
tater than the priority date claimed	*&" document member of the same patent family
Date of the actual completion of the international search	Date of mailing of the international search report
11 April 2000	09/05/2000
Name and mailing address of the ISA	Authorized officer
European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo ni, Fax: (+31-70) 340-3016	Fuchs, U

Form PCT/ISA/210 (second sheet) (July 1992)

Int. .tional Application No PCT/EP 99/09284

Category *	ation) DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages	In.
		Relevant to claim No.
	page 817; figures 2A-C page 818; figures 3A-F	
X	EMBL Database, Heidelberg, FRG Emest2 accession number AA700891 22 December 1997 Hillier, L. ET AL.: "zj40d01.s1 Soares fetal liver spleen 1NFLS S1 Homo sapiens cDNA clone 452737 3'" XP002135284 the whole document	6,7
X	EMBL Database, Heidelberg, FRG Emest6 accession number AI089668 19 August 1998 NCI/NINDS-CGAP: "qa10f06.x1 NCI_CGAP_Brn23 Homo sapiens cDNA clone IMAGE:1686371 3'" XP002135285 the whole document	6,8
x	BIRO, T. ET AL.: "Recent Advances in Understanding of Vanilloid Receptors: A Therapeutic Target for Treatment of Pain and Inflammation in Skin" JOURNAL OF INVESTIGATIVE DERMATOLOGY SYMPOSIUM PROCEEDINGS, vol. 2, no. 1, August 1997 (1997-08), pages 56-60, XP002075021	48,49
A	abstract page 57; table 1 page 58, column 1, line 8 -column 2, line 16	50,51
P,X	WO 99 37675 A (THE REGENTS OF THE UNIVERSITY OF CALIFORNIA) 29 July 1999 (1999-07-29)	1,2,4,6, 7,9,10, 12,14, 15,23, 24,26, 45,46, 48,50
	abstract page 1, line 1 -page 3, line 30 SEQ ID NOS: 33 and 34 page 100 -page 106	48,50
	page 58 -page 59; claims 1,24-6,8-14,19	
	-/	

Inte stional Application No
PCT/EP 99/09/284

C /Ca=1:=	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	PCT/EP 99/09284	
Category °			Relevant to claim No.
P,X	EP 0 943 683 A (SMITHKLINE BEECHAM PLC) 22 September 1999 (1999-09-22)		1,2,4,6, 7,9,10, 12,14, 15,23, 24,26, 45,46
	abstract page 2, line 1 - line 31 SEQ ID NOS: 1 and 2 page 14-16 page 36 -page 37; claims 1-14		
			·

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(a)

INTERNATIONAL SEARCH REPORT

...ernational application No.

PCT/EP 99/09284

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. X Claims Nos.: 27-45 because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically: See FURTHER INFORMATION sheet PCT/ISA/210
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims: it is covered by claims Nos.:
Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

International Application No. PCT/EP 99 09284

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box I.2

Claims Nos.: 27-45

Claims 27 - 45 refer to a compound which modulates human vanilloid receptor activity without giving a true technical characterization. Moreover, except two compounds already known in the prior art, no such compounds are defined in the application. In consequence, the scopes of said claims are ambigous and vague, and their subject matter is not sufficiently disclosed and supported (Art. 5 and 6 PCT).

No search can be carried out for such purely speculative claims whose wording is, in fact, a mere recitation of the result to be achieved.

The applicant's attention is drawn to the fact that claims, or parts of claims, relating to inventions in respect of which no international search report has been established need not be the subject of an international preliminary examination (Rule 66.1(e) PCT). The applicant is advised that the EPO policy when acting as an International Preliminary Examining Authority is normally not to carry out a preliminary examination on matter which has not been searched. This is the case irrespective of whether or not the claims are amended following receipt of the search report or during any Chapter II procedure.

Information on patent family members

Int. donal Application No PCT/EP 99/09284

Patent document cited in search report	rt	Publication date	!	Patent family member(s)	Publication date
WO 9937675	A	29-07-1999	AU AU WO	2466799 A 9115698 A 9909140 A	09-08-1999 08-03-1999 25-02-1999
EP 0943683	Α	22-09-1999	JP	11279196 A	12-10-1999

Form PCT/ISA/210 (patent family annex) (July 1992)

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My Williams